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(11) EP 1 415 996 A2

(12)

# **EUROPEAN PATENT APPLICATION**

(43) Date of publication: 06.05.2004 Bulletin 2004/19

(51) Int CI.7: **C07K 14/39**, C12N 9/12, C12N 1/15

(21) Application number: 03256895.8

(22) Date of filing: 30.10.2003

(84) Designated Contracting States:

AT BE BG CH CY CZ DE DK EE ES FI FR GB GR
HU IE IT LI LU MC NL PT RO SE SI SK TR
Designated Extension States:

AL LT LV MK

(30) Priority: 31.10.2002 JP 2002317736

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(54) Transformed cell with enhanced sensitivity to antifungal compound and use thereof

(57) The present invention provides a transformed cell in which a polynucleotide having a nucleotide sequence encoding an amino acid sequence of an osmosensing histidine kinase having no transmembrane region has been introduced in a functional form into a cell

deficient in at least one hybrid-sensor kinase, a method of assaying the antifungal activity of a test substance using the transformed cell, and a method of identifying an antifungal compound using the method, and the like.

#### Description

### BACKGROUND OF THE INVENTION

5 Field of the invention

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[0001] The present invention relates to a transformed cell with enhanced sensitivity to an antifungal compound and use thereof.

10 Description of the related art

[0002] It is known that, when a fungicide containing a dicarboxyimide antifungal compound, an "aromatic hydrocarbon antifungal compound" or a phenylpyrrole antifungal compound as an active ingredient is acted on a certain plant-pathogenic filamentous fungus, glycerol synthesis in a cell is stimulated in the fungus like as when undergoes high osmotic stress, and the fungus can not control an intracellular osmolarity, leading to death. From such the activity to the plant-pathogenic filamentous fungus, a protein in a signal transduction system which is involved in osmolarity response was predicted as a target protein of an antifungal compound contained in these fungicides as an active ingredient.

[0003] In Neurospora crassa exhibiting sensitivity to the aforementioned antifungal compound, an osmosensitive mutant os-1 was reported. This mutant os-1 exhibited resistance to the aforementioned antifungal compound and, by analysis of the mutant, an os-1 gene which is an osmosensing histidine kinase gene was isolated as a causative gene. A protein having an amino acid sequence encoded by a nucleotide sequence of this os-1 gene was a protein which has a structure of histidine kinase of a two-component regulatory system and, at the same time, has a characteristic region (hereinafter, referred to repeat sequence region in some cases) in which amino acid sequences composed of about 90 amino acids and having homology to each other are present repetitively about 6 times (see, for example, U. S.P. NO 5,939,306; Genebank accession U50263, U53189, AAB03698, AAB01979; Alex, A.L. et al., Proc. Natl. Red. Sci. USA 93:3416-3421; Schumacher, M.M. et al., Current Microbiology 34:340-347; Oshima, M. et al., Phytopathology 92 (1):75-80; Fijimura, M. et al., J. Pesticide Sci. 25:31-36). A gene having homology to the os-1 gene was also isolated fromplant-pathogenic filamentous fungus such as Botryotinia fuckeliana, Magnaporthe grisea, Fusarium solani and the like, and its nucleotide sequence and an amino acid sequence encoded by the gene are published. It is known that genes having homology with the os-1 gene are specifically present in filamentous fungus among eukaryotic organisms (see, for example, GeneBank accession AF396827, AF435964, AAL37947, AAL30826; Fujimura, M. et al., Pesticide Biochem. Physiol. 67:125-133; GeneBank accession AB041647, BAB40497).

## SUMMARY OF THE INVENTION

[0004] An object of the present invention is to provide a method of detecting the antifungal activity and a method of selecting an antifungal compound using the os-1 gene and a gene having homology with the gene.

[0005] Under such the circumstances, the present inventor intensively studied and, as a result, found a transformed cell with enhanced sensitivity to an antifungal compound, and found a method of detecting the antifungal activity using this transformed cell and a method of selecting an antifungal compound using this transformed cell, which resulted in completion of the present invention.

[0006] Thus, the present invention provides:

- 1. A transformed cell in which a polynucleotide having a nucleotide sequence encoding an amino acid sequence of an osmosensing histidine kinase having no transmembrane region is introduced in a functional form into a cell deficient in at least one hybrid-sensor kinase:
- 2. The transformed cell according to the above 1, the polynucleotide having a nucleotide sequence encoding an amino acid sequence of an osmosensing histidine kinase having no transmembrane region is a polynucleotide complementing the deficiency in hybrid-sensor kinase in the cell deficient in at least one hybrid-sensor kinase in which the polynucleotide is introduced;
- 3. The transformed cell according to the above 1, wherein the cell is a microorganism;
- 4. The transformed cell according to the above 3, wherein the microorganism is budding yeast;
- 5. The transformed cell according to the above 1, wherein the osmosensing histidine kinase having no transmembrane region is an osmosensing histidine kinase having no transmembrane region and having a mutation which confers resistance to any of a dicarboxyimide antifungal compound, an "aromatic hydrocarbon antifungal compound" and a phenylpyrrole antifungal compound to the cell;
- 6. The transformed cell according to the above 5, wherein the osmosensing histidine kinase having no transmem-

brane region is a histidine kinase having the amino acid sequence represented by SEQ ID NO: 13;

- 7. The transformed cell according to the above 1, wherein the osmosensing histidine kinase having no transmembrane region is an osmosensing histidine kinase which is derived from a plant-pathogenic filamentous fungus and has no transmembrane region;
- 8. The transformed cell according to the above 1, wherein the osmosensing histidine kinase having no transmembrane region is an osmosensing histidine kinase which is derived from Botryotinia fuckeliana, Magnaporthe grisea, Fusarium oxysporum, Mycospharella tritici, Thanatephorus cucumeris or Phytophthora infestans, and has no transmembrane region;
- 9. The transformed cell according to the abovel, wherein the osmosensing histidine kinase having no transmembrane region is an osmosensing histidine kinase having no transmembrane region which has an amino acid sequence represented by SEQ ID NO: 1, SEQ ID NO: 16, SEQ ID NO: 41, SEQ ID NO: 55, SEQ ID NO: 68 or SEQ ID NO: 90;
- 10. The transformed cell according to the above 1, wherein the nucleotide sequence encoding an amino acid sequence of the osmosensing histidine kinase having no transmembrane region is a nucleotide sequence represented by SEQ ID NO: 2, SEQ ID NO: 17, SEQ ID NO: 42, SEQ ID NO: 56 or SEQ ID NO: 69;
- 11. Amethod of assaying the antifungal activity of a substance, which comprises:

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- a first step of culturing the transformed cell as defined in the above 1 in the presence of a test substance; a second step of measuring an amount of intracellular signal transduction from the osmosensing histidine kinase having no transmembrane region expressed in the transformed cell cultured in the first step or an index value having the correlation therewith; and
- a third step of assessing the antifungal activity of the test substance based on a difference between an amount of intracellular signal transduction or an index value having the correlation therewith measured in the second step and a control;
- 12. The method of assaying according to the above 11, wherein the amount of intracellular signal transduction from the osmosensing histidine kinase having no transmembrane region or the index value having the correlation therewith is an amount of growth of the transformed cell;
- 13. A method of searching an antifungal compound, which comprises selecting an antifungal compound based on the antifungal activity assessed in the assaying method as defined in the above 11;
- 14. An antifungal compound selected by the searching method as defined in the above 13;
- 15. An osmosensing histidine kinase having no transmembrane region, wherein the kinase is derived from a plant-pathogenic filamentous fungus;
- 16. An osmosensing histidine kinase having no transmembrane region, which has an amino acid sequence selected from the group consisting of:
  - (a) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which has an amino acid sequence homology of 95% or more to the amino acid sequence represented by any of SEQ ID NOs: 41, 55 and 68;
  - (b) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Fusarium oxysporum-derived cDNA as a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 52 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 53 as primers;
  - (c) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Mycospharella tritici-derived cDNA as a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 64 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 65 as primers;
  - (d) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Thanapethorus cucumeris-derivedcD-NAas a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 85 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 86 as primers;
  - (e) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is derived from Phytophthora infestans and has the amino acid sequence represented by SEQ ID NO: 90;
  - (f) the amino acid sequence represented by SEQ ID NO: 41;
  - (g) the amino acid sequence represented by SEQ ID NO: 55; and
  - (h) the amino acid sequence represented by SEQ ID NO: 68.
- 17. An osmosensing histidine kinase having no transmembrane region, which has an amino acid sequence rep-

resented by SEQ ID NO: 41, SEQ ID NO: 55 or SEQ ID NO: 68;

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18. A polynucleotide having a nucleotide sequence encoding an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, wherein the kinase is derived from a plant-pathogenic filamentous fungus; 19. A polynucleotide having a nucleotide sequence encoding an amino acid sequence selected from the group consisting of:

- (a) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which has an amino acid sequence homology of 95% or more to the amino acid sequence represented by any of SEQ ID NOs: 41, 55 and 68;
- (b) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Fusarium oxysporum-derived cDNA as a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 52 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 53 as primers:
- (c) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Mycospharella tritici-derived cDNA as a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 64 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 65 as primers;
- (d) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Thanapethorus cucumeris-derived cDNA as a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 85 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 86 as primers;
- (e) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is derived from Phytophthora infestans and has the amino acid sequence represented by SEQ ID NO: 90;
- (f) the amino acid sequence represented by SEQ ID NO: 41;
- (g) the amino acid sequence represented by SEQ ID NO: 55; and
- (h) the amino acid sequence represented by SEQ ID NO: 68:
- 20. A polynucleotide having a nucleotide sequence represented by SEQ ID NO: 42, SEQ ID NO: 56 or SEQ ID NO:69;
- 21. A method of obtaining a polynucleotide having a nucleotide sequence encoding an amino acid sequence of osmosensing histidine kinase which is derived from a plant-pathogenic filamentous fungus and has no transmembrane region, which comprises a step of amplifying a desired polynucleotide by Polymerase Chain Reaction using an oligonucleotide having a nucleotide sequence represented by any of SEQ ID NOs: 30 to 40, 52, 53, 64, 65, 85 and 86 as primers, and a step of recovering the amplified desired polynucleotide; and
- 22. An oligonucleotide which comprises a nucleotide sequence represented by any of SEQ ID NOs: 30 to 40, 52, 53, 64, 65, 85 and 86.

## DETAILED DESCRIPTION OF THE PRESENT INVENTION

40 [0007] The present invention will be explained in detail below.

[0008] The "transformed cell in which a polynucleotide having a nucleotide sequence encoding an amino acid sequence of an osmosensing histidine kinase having no transmembrane region is introduced in a functional form into a cell deficient in at least one hybrid-sensor kinase" is obtained by introducing a polynucleotide having a nucleotide sequence encoding an amino acid sequence of an "osmosensing histidine kinase having no transmembrane region" in a functional form into a "cell deficient in at least one hybrid-sensor kinase" which is a host cell. Herein, "introduction of a polynucleotide in a functional form" means that the polynucleotide is introduced so as to complement the deficiency in hybrid-sensor kinase, in other words, that the polynucleotide is introduced in such a form that a phenotype of the cell caused by the deficiency in hybrid-sensor kinase revert to a phenotype without the deficiency in hybrid-sensor kinase. Specifically, for example, in the case of budding yeast (e.g. Saccharomyces cerevisiae), when SLN1 which is hybrid-sensor kinase is deleted, the SLN1-deficient yeast cell shows a phenotype that the cell can not grow under the normal growing condition. By introducing a polynucleotide having a nucleotide sequence encoding an amino acid sequence of SLN1 isolated from budding yeast into the SLN1-deficient cell so that SLN1 is expressed (e.g. operably linked to downstream of a promoter), the cell becomes possible to grow under the normal growing condition. The "cell deficient in at least one hybrid-sensor kinase" may be obtained, for example, by deleting at least one intrinsic hybrid-sensor kinase. First, hybrid-sensor kinase will be explained below.

(Two-component regulatory system and hybrid-sensor kinase)

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[0009] Two-component regulatory system is a signal transduction system which is widely used in prokaryotic organisms and, since this system is basically composed of two proteins called a sensor and a regulator, it is called two-component regulatory system. In a typical two-component regulatory system, a sensor is composed of an input region and a histidine kinase region, and a regulator is composed of a receiver region and an output region. When the input region senses an environmental stimulus, a histidine residue in an amino acid sequence in the histidine kinase region which is well conserved among organisms is phosphorylated or dephosphorylated. Herein, phosphorylation of the histidine residue is autophosphorylation utilizing ATP as a substrate. This phosphate group is transferred to an aspartic acid residue in an amino acid sequence in the receiver region in the regulator which is well conserved among organisms, and phosphorylation and dephosphorylation of the aspartic acid residue regulates the activity of the output region in the regulator. In the case of prokaryotic organisms, the output region is a transcription regulating factor in many cases although there are exceptions, and the regulator directly controls gene expression through the aforementioned phosphoryl transfer in response to stimuli sensed by the sensor.

[0010] A sensor takes a more complicated structure in some cases unlike the aforementioned typical structure. For example, in addition to a structure composed of an input region and a histidine kinase region, following this, the sensor contains a receiver region, which is observed in a regulator, on its C-terminal side in some cases. In this case, the phosphorylay system of a phosphate group becomes more complicated, and it is known that a phosphate is transferred from the sensor to a regulator called a response regulator via an intervening protein having a transmitter region called a phosphotransmitter. That is, when the input region of the sensor senses stimuli, phospate is transferred to mediate signal transduction from a histidine residue of the histidine kinase region in the same molecule to an aspartic acid residue of the receiver region in the same molecule, then, to a histidine residue of the phosphotransmitter, finally, to an aspartic acid residue of the receiver region in a response regulator. Like this, two-component regulatory system is associated with three proteins in some cases. Such the sensor involved in signal transduction system through phosphoryl transfer composed of three proteins and having the aforementioned structural characteristic is referred to as "hybrid-sensor kinase". Hybrid-sensor kinase is found not only in a prokaryotic organism but also in an eukaryoticmicroorganism such as yeast, a plant and the like, and is involved in response to a variety of stimuli or stresses.

[0011] Herein, an input region of a hybrid-sensor kinase is a region present at the N-terminal of the kinase, and have a transmembrane region in many cases. The transmembrane region can be revealed by a structure prediction analysis using a structure prediction software, for example, TMpred program [K. Hofmann & W. Stoffel, Biol. Chem. Hoppe-Seyler, 374, 166 (1993)] which is available, for example, from http://www.ch.embnet.org/software/TMPRED\_form.html. A histidine kinase region of a hybrid-sensor kinase is, for example, a region following the C-terminal of the input region, and is a region characterized in that it has five conserved motifs common to general histidine kinases as described in Parkinson, J.S. & Kofoid, E.C. (1989) Annual Review of Genetics 23:311-336, Stock, J.B. et. al. (1989) Microbiological Reviews 53 (4):450-490. For example, in the hybrid-sensor kinase SLN1 of budding yeast, a histidine kinase region following the C-terminal of the histidine kinase region, and is a region characterized in that it has three conservedmotifs common to general histidine kinases as described in Parkinson, J.S. & Kofoid, E.C. Annual Review of Genetics 23: 311-336(1989), Stock, J.B. et. al. (1989) Microbiological Reviews 53 (4): 450-490. For example, in the hybrid-sensor kinase SLN1 of budding yeast, a receiver region is-the region from amino acid residues 1088 to 1197.

[0012] As a signal transduction systemafter a response regulator, in addition to a simple system in which an output region of a regulator is a transcription regulating factor as described above, as a more complicated system, there is known a system in which a signal is transmitted to a transcription regulating factor participating in control of gene expression, via MAP kinase cascade which is associated with various controls in a cell.

[0013] Specific examples of a hybrid-sensor kinase and a signal transduction systemwhich involves the hybrid-sensor kinase will be explained below.

(Hybrid-sensor kinase of budding yeast)

[0014] In budding yeast (Saccharomyces cerrevisiae), the hybrid-sensor kinase SLN1 is utilized for signal transduction relating to osmolarity response. The SLN1 is a sole histidine kinase found in budding yeast. SLN1 is an osmosensing histidine kinase having a transmembrane region in its input region, and mediates a phosphoryl transfer signal to the response regulator SSK1 via the phosphotransmitter YPD1. Downstream of the signal transduction, MAP kinase cascade composed of three kinases SSK2(MAPKKK), PBS2 (MAPKK) and HOG1 (MAPK) lies to regulate expression of genes involved in osmolarity adaptation such as glycerol biosynthesis and the like. The output region of the response regulator SSK1 has an activity of phosphorylating SSK2. The SSK1 is negatively controlled by phophorylation of an aspartic acid residue in its receiver region, the phosphorylating activity of whose output region is inhibited. Specifically, at a normal osmolarity, a histidine residue in the histidine kinase region of SLN1 is autophosphorylated, and the phos-

phate is subsequently transferred to an aspartic acid residue of the receiver region in the same molecule, then, to a histidine residue of YPD1, finally, to an aspartic acid residue in the receiver region of SSK1. By phosphorylation of an aspartic acid residue in the receiver region of SSK1, the phosphorylating activity of the output region of SSK1 is suppressed, and the phosphate is not transferred to a MAP kinase cascade composed of SSK2, PBS2 and HOG1, and then expression of genes involved in osmolarity adaptation such as glycerol biosynthesis and the like are not induced. On the other hand, under a condition of high osmolarity, since autophosphorylation of a histidine residue of the histidine kinase region is inhibited in SLN1, the MAP kinase cascade composed of SSK2, PBS2 and HOG1 is activated, and then expression of genes involved in osmolarity adaptation such as glycerol biosynthesis and the like is induced (Maeda, T. et. al. (1994) Nature 369:242-245).

(Hybrid-sensor kinase of fission yeast)

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[0015] In fission yeast (Scchizosaccharomyces pombe), three kinds of hybrid-sensor kinases PHK1 (MAK2), PHK2 (MAK3) and PHK3 (MAK1) participate in regulation of cell cycle progression [G (2) to M phase transition] and oxidative stress response. In a fission yeast, there is no histidine kinase other than PHK1, PHK2 and PHK3. PHK1 and PHK2 are histidine kinases responsive to oxidative stress such as hydrogen peroxide and the like (Buck, V. et. al., Mol. Biol. Cell 12:407-419). Three kinds of hybrid-sensor kinases PHK1, PHK2 and PHK3 metiate a phosporyl transfer signal to the response regulator MCS4 via the phosphotransmitter SPY1 (MPR1). Downstream of this signal transduction, a MAP kinase cascade composed of three kinases WAK1 (MAPKKK), WIS1 (MAPKK) and STY1 (MAPK) lies to regulate expression of genes involved in regulation of cell cycle progression and oxidative stress response. The output region of the response regulator MCS4 has an activity of phosphorylating WAK1. The MCS4 is negatively controlled by phosphorylation of an aspartic acid residue in its receiver region, the phosphorylating activity of whose output region is inhibited. Specifically, under a normal condition, each of histidine residues in the histidine kinase regions of PHK1 to PHK3 is autophosphorylated, and the phosphates are transferred to each of aspartic acid residues of receiver regions in the same molecule, then, to a histidine residue of SPY, finally, to an aspartic acid residue in the receiver region of MCS4. By phosphorylation of an aspartic acid residue in the receiver region of MCS4, the phophorylating activity of the output region of MCS4 is suppressed, and the phosphate is not transferred to a MAP kinase cascade composed of WAK1, WIS1 and STY1, and then expression of genes involved in regulation of cell cycle progression and stress response are not induced. On the other hand, under a stress condition, autophosphorylation of each of histidine residues of the histidine kinase regions in PHK1 to PHK3 is inhibited, a MAP kinase cascade composed of WAK1, WIS1 and STY1 is activated, and expression of genes involved in control of cell cycle progression and oxidative stress response are induced. As a result, it is observed such a phenotype that G (2) to M phase transition in cell cycle progression of the fission yeast is promoted, and that a dividing cell length becomes remarkably shorter than usual (Aoyama, K. et. al. (2001) Boisci. Biotechnol. Biochem. 65:2347-2352).

-(Hybrid-sensor kinase of bacterium)

[0016] In a prokaryotic organism Escherichia coli, the hybrid-sensor kinase RcsC participates in control of expression of the cps operon involved in capsular polysaccharide synthesis. RcsC is a histidine kinase having a transmembrane region, and it is known that it mediates a phosphoryl transfer signal to the response regulator RcsB via the phosphotransmitter YojN. The output region of RcsB has an activity of inducing transcription of the cps operon. Specifically, under a normal condition, a histidine residue in the histidine kinase region of RcsC is autophosphorylated, and the phosphate is transferred to an aspartic acid residue of the receiver region in the same molecule, then, to a histidine residue of YojN, finally, to an aspartic acid residue in the receiver region of RcsB. By phosphorylation of an aspartic acid residue in the receiver region of RcsB. By phosphorylation of an aspartic acid residue in the receiver region of RcsB, the cps operon transcription inducing activity of the output region of RcsB is suppressed, and expression of genes involved in capsular polysaccharide synthesis are not induced. On the other hand, under a condition of high osmolarity, in RcsC, autophosphorylation of a histidine residue in the histidine kinase region is inhibited, the cps operon transcription inducing activity of the output region of RcsB is activated, and expression of genes involved in capsular polysaccharide synthesis are induced (Clarke, D. J. et. al. (2002) J. Bactriol. 184: 1204-1208).

[0017] A bioluminescent marine microorganism Vibrio harveyi emits fluorescent light generated in luciferase reaction depending on its own cell density. Hybrid-sensor kinases LuxN and LuxQ parcipite in control of expression of a gene involved in this cell density-responsive bioluminescence. LuxN and LuxQ are histidine kinases each having a transmembrane region. To sense its own cell density, V. harveyi produces and secrets two kinds of substances (Al-1, Al-2) called autoinducer. Al-1 is sensed by LuxN and Al-2 is sensed by LuxQ to convey cell-density information. LuxN and LuxQ mediate phosphoryl transfer signals to the response regulator LuxO via the phosphotransmitter LuxU. The output region of LuxO has an activity of inducing transcription of the luciferase operon. To specifically explain by referring to LuxN, when a cell density is low, since Al-1 in the environment is at low level and is not sensed by the input region of

LuxN, a histidine residue in the histidine kinase region of LuxN is autophosophorylated. The phosphate is transferred to an aspartic acid residue of the receiver region in the samemolecule, then, to a histidine residue of LuxU, finally, to an aspartic acid residue in the receiver region of LuxO. By phosphorylation of an aspartic acid residue in the receiver region of LuxO, the luciferase operon transcription inducing activity of the output region of LuxO is suppressed, and expression of genea involved in bioluminescence are not induced. On the other hand, under a high cell density condition; since Al-1 in environment is at high level and is sensed by the input region of LuxN, autophosphorylation of a histidine residue of the histidine kinase region is inhibited in LuxN, the luciferase operon transcription inducing activity of the output region of LuxO is activated, andbioluminescence is induced (Freeman, J.A. et.al. (2000) Mol. Microbiol. 35:139-149).

(Hybrid-sensor kinase of plant)

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[0018] In a higher plant Arabadopsis thaliana, receptor proteins CRE1, AHK2 and AHK3 for a plant hormone cytokinin are hybrid-sensor kinases. Receptor proteins CRE1, AHK2 and AHK3 are all cytokinin-sensitive histidine kinase having a transmembrane region (Inoue, T. et. al.(2001) Nature 409:1060-1063). CRE1 mediates a phosphoryl transfer signal to response regulators ARR1, ARR2 and ARR10 via phosphotransmitters AHP1 and AHP2. It is considered that output regions of ARR1, ARR2 and ARR10 have an activity of inducing transcription of cytokinin-inducing genes ARR4 to ARR7. Specifically, in the presence of cytokinin, a histidine residue in the histidine kinase region of CRE1 is autophosphorylated, and the phosphate is transferred to an aspartic acid residue of the receiver region in the same molecule, then, to histidine residues of AHP1 and AHP2, finally, to aspartic acid residues in receiver regions of ARR1, ARR2 and ARR10. By phosphorylation of aspartic acid residues in receiver regions of ARR1, and ARR10, a gene transcription inducing activity of output regions of ARR1, ARR2 and ARR10 are promoted, and expression of cytokinin-responsive genes ARR4 to 7 is induced (Hwang, I. & Sheen J. (2001) Nature 413:383-389).

(Cell deficient in at least one hybrid-sensor kinase)

[0019] "The cell deficient in at least one hybrid-sensor kinase" means a cell in which function of at least one intrinsic hybrid-sensor kinase is lost. Examples of the cell include a cell in which production of at least one intrinsic hybrid-sensor kinase is deleted, suppressed or inhibited, a cell in which activity of at least one intrinsic hybrid-sensor kinase is deleted, suppressed or inhibited, and the like. More specific examples include budding yeast deficient in SLN1, fission yeast deficient in all of three of PHK1, PHK2 and PHK3, Escherichia coli deficient in RcsC, V. harveyi deficient in LuxN, Arabidopsis thaliana deficient in CRE1, and the like.

[0020] In order to prepare the "cell deficient in at least one hybrid-sensor kinase", for example, deletion, addition, substitution or the like of one or more nucleotides are introduced into the whole or a part of a promoter region or a coding region of a gene encoding hybrid-sensor kinase to be deleted. Specifically, for example, the SLN1-deficient-budding yeast strain TM182 can be prepared by the method described in Maeda, T. et. al. (1994) Nature 369:242-245, the PHK1, PHK2 and PHK3-deficient fission yeast strain Kl011 can be prepared by the method described in Aoyama, K. et. al. (2001) Boisci. Biotechnol. Biochem. 65:2347-2352. In addition, the RcsC-deficient Escherichia coli strain SRC122 can be prepared by the method described in Suzuki, T., et. al. (2001) Plant Cell Physiol. 42:107-113, and the LuxN-deficient V. harveyi strain BNL63 can be prepared by the method described in Freeman, J.A. et. al. (2000) Mol. Micobiol. 35:139-149. For preparing a CRE1-deficient Arabidopsis thaliana, for example, a clone defective in cytokine response is selected from clones obtained by mutagenesis of Arabidopsis thaliana according to the method described in Inoue, T. et. al. (2001) Nature 409:1060-1063. Genomic CRE1 gene fragment is amplified by PCR using a primer designed based on the nucleotide sequence of the genomic CRE1 gene listed in Genebank accession AB049934 and using a genomic DNA of the selected clone as a template, and its nucleotide sequence is confirmed, whereby, a CRE1-deficient clone which can not express CRE1 can be selected.

[0021] Alternatively, a cell deficient in unknown hybrid-sensor kinase besides the aforementioned kinases may be also prepared, for example, by isolating a hybrid-sensor kinase gene from a desired cell, and deleting the gene harbored by the cell by homologous recombination using the gene. For isolating a hybrid-sensor kinase gene of a desired cell, the structural characteristic of hybrid-sensor kinases can be utilized. For example, amino acid sequences around the histidine residue tobe autophosphorylated are conserved among histidine kinase regions and amino acid sequences around the aspartic acid residue to which a phosphate to be transferred from the histidine residue are conserved among receiver regions. Then, a hybrid-sensor kinase gene of a desired cell can be isolated by a polymerase chain reaction (hereinafter, referred to as PCR) using an oligonucleotide designed based on a nucleotide sequence encoding the aforementioned conserved amino acid sequences as a primer, or a hybridization method using an oligonucleotide having a nucleotide sequence encoding the aforementioned conserved amino acid sequences as a prove. By examining whether or not the aforementioned structural characteristic is possessed based on an amino acid sequence deduced from a nucleotide sequence of the isolated gene, it can be confirmed that the isolated gene is a gene having a nucleotide

sequence encoding an amino acid sequence of a hybrid-sensor kinase. A specific example is a PCRmethod described in Srilantha, T. et. al. (1998) Microbiology 144:2715-2729. For PCR and hybridization, for example, the experimental conditions using upon isolation of the "polynucleotide having a nucleotide sequence encoding an amino acid sequence of osmosensing hisitidine kinase having no transmembrane region" described later may be used.

[0022] Alternatively, a hybrid-sensor kinase gene may be also isolated using, as an index, the functional complementation in budding yeast in which expression of SLN1 is conditionally suppressed, for example, according to the method described in Nagahashi, S. et. al. (1998) Microbiology 144:425-432.

(Osmosensing histidine kinase having no transmembrane region)

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[0023] Then, the "osmosensing histidine kinase having no transmembrane region" to be introduced into the aforementioned "cell deficient in at least one hybrid-sensor kinase" in a functional form will be explained.

[0024] In filamentous fungus, a histidine kinase having a structure similar to that of the aforementioned hybrid-sensor kinase is isolated. The histidine kinase has a histidine kinase region and a receiver region which are observed in hybrid-sensor kinases, and has no transmembrane region, which is observed in many hybrid-sensor kinases, in its input region, and further has a characteristic structure in which amino acid sequences composed of about 90 amino acids having the amino acid sequence homology to each other are present repeatedly about six times, in place of the transmembrane region. Although a signal transduction pathway from this histidine kinase has not been completely clarified, it is known that the signal transduction participates in osmolarity response.

[0025] In the present invention, "homology" refers to identity of sequences between two genes or two proteins. The "homology" is determined by comparing two sequences aligned in the optimal state, over a region of a sequence of a subject to be compared. Herein, in optimal alignment of nucleotide sequences or amino acid sequences to be compared, addition or deletion (e.g. gap etc.) may be allowable. Such the "homology" can be calculated by homology analysis with making alignment using a program of FASTA [Pearson & Lipman, Proc. Natl. Acad. Sci. USA, 4, 2444-2448 (1998)], BLAST [Altschul et. al. Journal of Molecular Biology, 215, 403-410 (1990)], CLUSTAL W [Thompson, Higgins & Gibson, Nucleic Acid Research, 22, 9673-4680 (1994a)] and the like. The above programs are available to the public, for example, inhomepage (http://www.ddbj.nig.ac.jp) of DNADataBankof Japan [international DNA Data Bank managed in Center for Information Biology and DNA Data Bank of Japan (CIB/DDBJ)). Alternatively, the "homology" may be also obtained by using commercially available sequence analysis software. Specifically, the homology can be calculated, for example, by performing homology analysis with making alignment by the Lipman-Pearson method [Lipman, D. J. and Pearson, W. R., Science, 227, 1435-1441, (1985)] using GENETYX-WIN Ver.5 (manufactured by Software Development Co. Ltd.)

[0026] Herein, as the "structure in which amino acid sequences composed of about 90 amino acids having the amino acid sequence homology to each other are repeatedly present about six times", for example, there is a repeat sequence region described in Alex, L.A. et. al. (1996) Proc. Natl. Acad. Sci. USA 93:3416-3421, Ochiai, N. et. al. (2001) Pest Manag. Sci. 57:437-442, Oshima, M. et. al. (2002) Phytopathology 92:75-80 and the like, and such the structure is present at the N-terminal region of the hisitide kinase. The "amino acid sequences composed of about 90 amino acids are repeatedly present about six times" include an amino acid sequence motif composed of about 90 amino acids is repeated five times followed by a sixth truncated repeat sequence (5.7 times repeat), an amino acid sequence motif composed of about 90 amino acids is repeated six times followed by a seventh truncated repeat sequence (6.7 times repeat), and the like. Specifically, in amino acid sequence of a histidine kinase of the present invention, examples of the "a region in which amino acid sequences composed of about 90 amino acids having the amino acid sequence homology to each other are present repeatedly about six times" include a region from amino acid residues 190 to 707 in an amino acid sequence represented by SEQ ID NO: 1 (5.7 times repeat), a region from amino acid residues 189 to 706 in an amino acid sequence represented by SEQ ID NO: 16 (5.7 times repeat), a region from amino acid residues 176 to 693 in an amino acid sequence represented by SEQ ID NO: 41 (5.7 times repeat), a region from amino acid residues 192 to 709 in an amino acid sequence represented by SEQ ID NO: 55 (5.7 times repeat), and a region from amino acid residues 299 to 911 in an amino acid sequence represented by SEQ ID NO: 68 (6.7 times repeat), and the

[0027] The "osmosensing histidine kinase having no transmembrane region" is the aforementioned histidine kinase characteristic in filamentous fungus, and refers to a osmosensing protein having a repeat sequence region of amino acid sequences composed of about 90 amino acids having the amino acid sequence homology to each other, a histidine kinase region and a receiver region, and having no transmembrane region.

[0028] In order to confirm that a protein has the function of osmosensing histidine kinase, enhancement of the sensitivity of a cell to osmolarity stress may be confirmed when the protein (histidine kinase) is deleted from the cell. Alternatively, it may be also confirmed that a protein (histidine kinase) is osmosensing histidine kinase, by confirming that expression of the protein in an osmosensing hybrid-sensor kinase SLN1-deficient budding yeast cell results in a functional complementation of the SLN1 and the budding yeast cell capable of growing.

[0029] Among filamentous fungi, mainly, in Neurospora crassa which is a model organism of filamentous fungus, a plant pathogenic filamentous fungus which is a pathogenic microorganism, ahostofwhichisaplant, or the like, the presence of the "osmosensing histidine kinase having no transmembrane region" is reported.

[0030] Examples of the "osmosensing histidine kinase having no transmembrane region" of the present invention include an osmosensing histidine kinase having no transmembrane region, which has an amino acid sequence selected from the group consisting of:

- (a) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which has an amino acid sequence homology of 95% or more to the amino acid sequence represented by any of SEQ ID NOs: 41, 55 and 68;
- (b) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Fusarium oxysporum-derived cDNA as a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 52 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 53 as primers;
- (c) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Mycospharella tritici-derived cDNA as a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 64 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 65 as primers;
- (d) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Thanapethorus cucumeris-derived cDNAas a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 85 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 86 as primers;
- (e) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is derived from Phytophthora infestans and has the amino acid sequence represented by SEQ ID NO: 90;
- (f) the amino acid sequence represented by SEQ ID NO: 41;

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- (g) the amino acid sequence represented by SEQ ID NO: 55; and
- (h) the amino acid sequence represented by SEQ ID NO: 68.

[0031] A preferred amino acid sequence homology in the above (a) may for example be about 95%, or higher such as about 98%. The difference from the amino acid sequence represented by any of SEQ ID: 41, 55 and 68 observed in the amino acid sequence of the above (a) may for example be a variation such as the deletion, substitution and addition of amino acids. Such a variation includes a variation which can artificially be introduced by means of a site-directed mutagenesis method or a mutagenic treatment as well as a polymorphic variation which occurs naturally such as a difference in an amino acid sequence resulting from the difference by the species or strains from which the protein is derived. As the site-directed mutagenesis method, for example, there is mentioned the method which utilizes amber mutations (capped duplex method, Nucleic Acids Res., 12, 9441-9456 (1984)), the method by PCR utilizing primers for introducing a mutation and the like.

[0032] At least one, specifically one to several (herein"several" means about 2 to about 10), or more amino acid residues may be varied in the above variations. The amino acid residues may be varied in any numbers as far as the effect of the present invention can be observed.

[0033] Of the deletion, addition, and substitution, the substitution is particularly preferred in the amino acid variation. Amino acids that are similar to each other in hydrophobicity, charge, pK, stereo-structural characteristic, or the like are more preferably replaced with each other. For example, such substitutable amino acids are in each of the following groups: 1) glycine and alanine; 2) valine, isoleucine, and leucine; 3) aspartic acid, glutamic acid, asparagine, and glutamine; 4) serine and threonine; 5) lysine and arginine; and 6) phenylalanine and tyrosine.

[0034] The "osmosensing histidine kinase having no transmembrane region" will be further explained with the specific examples shown below.

(Osmosensing histidine kinase having no transmembrane region of Neurospora crassa)

[0035] A protein OS-1 encoded by an os-1 gene isolated from an osmosensing mutant os-1 of Neurospora crassa can be mentioned as the "osmosensing histidine kinase having no transmembrane region" (Schumacher, M. M. et. al. (1997) Current Microbiol. 34:340-347, Alex, L. A. et. al. (1996) Proc. Natl. Acad. Sci. USA 93:3416-3421). Amino acid sequences of OS-1 and nucleotide sequences of the os-1 gene are published (amino acid sequence: Genebank accession AAB03698, AAB01979, nucleotide sequence: Genebank accession U50263, U53189), and utility of OS-1 and os-1 gene in screening system for antifungal compounds is described in US 5, 939, 306. Since Neurospora crassa mutant os-1 has the higher sensitivity to high osmolarity stress than that of a wild strain, it has been found that OS-1 is an osmosensing histidine kinase involved in osmolarity adaptation in Neurospora crassa.

[0036] It is known that OS-1 has the aforementioned structural characteristic based on its amino acid sequence. In addition, it is known that Neurospora crassa mutant os-1 has the resistance to fungicides containing, as an active ingredient, a dicarboxyimide antifungal compound, an "aromatic hydrocarbon antifungal compound" or a phenylpyrrole antifungal compound.

[0037] Further, a gene mutation which leads to an amino acid substitution in a characteristic repeat sequence region of OS-1 was observed in the os-1 mutant gene isolated from Neurospora crassa mutant exhibiting the resistance to a fungicide containing a dicarboxyimide antifungal compound as an active ingredient (Miller, T. K. et. al. (2002) Fungl Gen. Biol. 35:147-155). From the foregoing, it is predicted that an antifungal compound contained as an effective ingredient in the aforementioned fungicide targets OS-1 of Neurospora crassa.

(Osmosensing histidine kinase having no transmembrane region of Botryotinia fuckeliana)

[0038] Examples of the "osmosensing histidine kinase having no transmembrane region" include BcOS-1 of Botryotinia fuckeliana. The BcOS-1 gene was isolated as a gene homologous to Neurospora crassa OS-1 gene, and nucleotide sequencez and amino acid sequences are published (nucleotide sequence: GeneBank accession AF396287,
AF435964, amino acid sequence: GeneBank accession AAL37947, AAL30826). It is known that BcOS-1 has the aforementioned structural characteristic based on its amino acid sequence. In addition, in the BcOS-1 gene isolated from
a Botryotinia fuckeliana strain resistant to a fungicide containing a dicarboxyimide antifungal compound as an active
ingredient, a mutation which leads to amino acid substitution in the characteristic repeat sequence region of BcOS-1
was observed, as in the OS-1 gene isolated from a Neurospora crassa strain resistant to a fungicide containing a
dicarboxyimide antifungal compound as an active ingredient. Further, since an antifungal compound-resistant mutant
deficient in the BcOS-1 has the higher osmolarity sensitivity than that of a wild strain, it is known that BcOS-1 is osmosensing histidine kinase (Oshima, M. et. al. (2002) Phypotathology 92:75-80).

[0039] More specifically, examples of BcOS-1 include BcOS-1 having an amino acid sequence represented by SEQ ID NO: J. which was isolated from Be-16 strain described in Example. (Osmosensing histidine kinase having no transmembrane region of Magnaporthe grisea)

[0040] Example of the "osmosensing histidine kinase having no transmembrane region" include HIKI of Magnaporthe grisea. The hik1 gene is a gene homologous to Neurospora crass os-1 gene, and a nucleotide sequence and an amino acid sequence are published (nucleotide sequence: Genebank accession AB041647, amino acid sequence: GeneBank accession BAH40947). It is known that HIK1 has the aforementioned structural characteristics such as lack of the transmembrane region based on its amino acid sequence. In addition, it is observed that Magnaporthe grisea deficient in the hik1 gene has the higher osmolarity sensitivity than that of a wild strain, demonstrating that HIK1 is an osmosensing histidine kinase

(hppt://www.sci.saitama-u.ac.jp/seitai/iden/Japanese/Abst Symp3. html).

[0041] More specifically, examples of HIK1 include HIK1 having an amino acid sequence represented by SEQ ID
 NO: 16 which was isolated from the P-37 strain described in Example.

(Definition of filamentous fungus and yeast)

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[0042] In the present invention, the "filamentous fungus" means fungi other than fungi which can be classified as yeast, among fungi consisting of Myxomycota and Eumycota, described in "Revised Edition, Classification and Identification of Microorganisms (Volume 1), edited by Takeharu HASEGAWA, Society Publishing Center, 1984 (ISDN 4-7622-7399-6)". Examples of filamentous fungus classified in Myxomycota include Plasmodiophora brassicae belonging to Plasmodiophoromycetes. In addition, examples of filamentous fungus which is classified in Eumycota include Phytophthora infestans belonging to Mastigomycotina, Rhizopus stolonifer and Rhizopus oryzae belonging to Zygomycotina, Neurospora crassa, Mycospharella tritici, Erysiphe graminis, Linocarpon cariceti, Cochliobolus miyabeanus, Botrytinia fuckeliana and Magnaporthe grisea belonging to Ascomycotina, Ustilago maydis, Puccinia recondite and Thanatephorus cucumeris belonging to Basidiomycotina, Cladosporium fulvum, Alternalia kikuchiana and Fusarium oxysporum belonging to Deuteromycotina, and the like.

[0043] In addition, yeast means fungi in which they are grown mainly by budding, a single cell generation is long, a colony formed by growth of a single cell does not become hairy, but becomes white bright paste-like as described in "Revised Edition, Classification and Identification of Microorganisms (Volume 1), edited by Takeharu HASEGAWA, Society Publishing Center, 1984 (ISBN 4-7622-7399-6)". Examples thereof include Saccharomyces cerevisiae belonging to genus Saccharomyces, Schizosaccharomyces pombe belonging to genus Schizosaccharomyces, Phichia burtonii belonging to genus Phichia, Candida albicans belonging to genus Candida, and the like.

(Osmosensing histidine kinase having mutation which confers resistance to any of dicarboxyimide antifungal compound, aromatic hydrocarbon antifungal compound and phenylpyrrole antifungal compound, and having no transmembrane region)

[0044] As a specific example of the "osmosensing histidine kinase having no transmembrane region", there can also be exemplified "osmosensing histidine kinase having no transmembrane region" having mutation which confers resistance to any of a dicarboxyimide antifungal compound, an "aromatic hydrocarbon antifungal compound" and a phenylpyrrole antifungal compound. Specifically, there can be exemplified BcOS-1 having an amino acid sequence represented by SEQ ID NO: 13 which is described in Example.

[0045] Herein, the dicarboxyimide antifungal compound is a generic name of antifungal compounds having dicarboxyimide as a basic structure, and examples thereof include antifungal compounds described in Modern Selective Fungicide-Properties, Applications, Mechanism of Action-2<sup>nd</sup> revised and enlarged edition Lyr, H. ed. Gustav Fisher Verlag, New York, USA ISBN 3-334-60455-1 Chapter 6, p99-118. Specifically, there are a compound having a structure represented by the chemical formula (1) (Procymidone: hereinafter, referred to as Compound (1) in some cases), a compound having a structure represented by the chemical formula (2) (Iprodione: hereinafter, referred to as Compound (2) in some cases), a compound having a structure represented by the chemical formula (3) (Vinclozolin: hereinafter, referred to as Compound (3) in some cases) and the like. The "aromatic hydrocarbon antifungal compound" is a generic name of antifungal compounds having a benzene ring as a basic structure, and examples thereof include antifungal compounds described in Modern Selective Fungicide-Properties, Applications, Mechanism of Action-2nd revised and enlarged edition Lyr, H. ed. Gustav Fisher Verlag, New York, USA ISBN 3-334-60455-1 Chapter 5, p75-98. Specifically, there are a compound having a structure represented by the chemical formula (4) (Quintozene: hereinafter, referred to as Compound (4) in some cases), a compound having a structure represented by the chemical formula (5) (Tolclofosmethyl: hereinafter, referred to as Compound (5) in some cases). In addition, the phenylpyrrole antifungal compound is a generic name of antifungal compounds having phenylpymole as a basic structure, and examples thereof include antifungal compounds described in Modern Selective Fungicide-Properties, Applications, Mechanism of Action-2nd revised and enlarged edition Lyr, H. ed. Gustav Fisher Verlag, New York, USA ISBN 3-334-60455-1 Chapter 19, p405-407. Specifically, there are a compound having a structure represented by the chemical formula (6) (Fludioxonil: hereinafter, referred to as Compound (6) in some cases), a compound having a structure represented by the chemical formula (7) (Fenpiclonil: hereinafter, referred to as Compound (7) in some cases) and the like.

[0046] Chemical formulas of the aforementioned dicarboxyimide antifungal compound, "aromatic hydrocarbon antifungal compound" and phenylpyrrole antifungal compounds are shown below.

(1) Compound having a structure represented by the chemical formula (1) (Compound (1))

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# Chemical formula (1)

(2) Compound having a structure represented by the chemical formula (2) (Compound (2))

# Chemical formula (2)

(3) Compound having a structure represented by the chemical formula (3) (Compound (3))

# Chemical formula (3)

5 CI CH<sub>3</sub> CH<sub>3</sub>

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(4) Compound having a structure represented by the chemical formula (4) (Compound (4))

# Chemical formula (4)

C | 01

(5) Compound having a structure represented by the chemical formula (5) (Compound (5))

# Chemical formula (5)

сну осну

(6) Compound having a structure represented by the chemical formula (6) (Compound (6))

# Chemical formula (6)

NG NH

# (7) Compound having a structure represented by the chemical formula (7) (Compound (7))

# Chemical formula (7)

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[0047] The "mutation which confers resistance to any of a dicarboxyimide antifungal compound, an aromatic hydrocarbon antifungal compound and a phenylpyrrole antifungal compound" indicates a mutation which can be found in the "osmosensing histidine kinase having no transmembrane region" produced by a filamentous fungus mutant having resistance to any of a dicarboxyimide antifungal compound, an "aromatic hydrocarbon antifungal compound" and a phenylpyrrole antifungal compound, that is, substitution, addition or deletion of one or more amino acids which confer resistance to a dicarboxyimide antifungal compound, an "aromatic hydrocarbon antifungal compound" and a phenylpyrrole antifungal compound, provided that mutation by which the "osmosensing histidine kinase having no transmembrane region" becomes not to function as histidine kinase is eliminated. Herein, a mutant of filamentous fungus having resistance to any of a dicarboxyimide antifungal compound, an "aromatic hydrocarbon antifungal compound" and a phenylpyrrole antifungal compound may be filamentous fungus isolated from the nature to which any of a dicarboxyimide antifungal compound, an "aromatic hydrocarbon antifungal compound" and a phenylpyrrole antifungal compound was applied, or may be resistance-acquired filamentous fungus selected by artificially culturing filamentous fungus in the presence of a dicarboxyimide antifungal compound, an "aromatic hydrocarbon antifungal compound" or phenylpyrrole antifungal compound.

[0048] Specifically, in BcOS-1 in the "osmosensing histidine kinase having no transmembrane region" of Botryotinia fuckeliana, amino acid-substitution I365S which confers resistance to a dicarboxyimide antifungal compound is reported in Oshima, M. etal. (2002) Phytopathology 92: 75-80 (herein, "I365S" means that isoleucine at amino acid residue 365 is substituted with serine. Hereinafter, amino acid substitution is described similarly). As an amino acid substitution which confers resistance to a dicarboxyimide antifungal compound in OS-1 which is the "osmosensing histidine kinase having no transmembrane region" of Neuorspora crassa, T368P, Q388S, E418E, L459M, A578V, G580R, I582M, -M639V, A578V, G580G and L625P are reported and, as an amino acid deletion, 680K is reported in Miller, T.K. et al. (2002) Fungal Gen. Biol. 35:147-155 (hereinafter, 680Kmeans that lysine at amino acid residue 680 is deleted. Hereinafter, amino acid deletion is described similarly). In addition, amino acid substitution which confers resistance to a phenylpyrrole antifungal compound in the OS-1 of Neurospora crassa, A578V, G580R and L625P are reported in

Ochiai, N. et al. (2001) Pest Management Sci. 57:437-442.

[0049] Besides the aforementioned resistance mutation, resistance mutation may be found by analyzing an amino acid sequence of the "osmosensing hystidine kinase having no transmembrane region" isolated from a mutant filamentous fungus having resistance to anyof a dicarboxyimide antifungal compound, an "aromatic hydrocarbon antifungal compound" and a phenypyrrole antifungal compound, and comparing with an amino acid sequence of the protein in a sensitive wild strain.

(Preparation of transformed cell in which a polynucleotide having a nucleotide sequence encoding an amino acid sequence of osmosensing histidine kinase having no transmembrane region is introduced in a functional form into a cell deficient in at least one hybrid-sensor kinase)

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[0050] The transformed cell in which a polynucleotide having a nucleotide sequence encoding an amino acid sequence of osmosensing histidine kinase having no transmembrane region (hereinafter, referred to as present histidine kinase in some cases) is introduced in functional form, can be obtained by introducing a "polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present hystidine kinase" or the like into a "cell deficient in at least one hybrid-sensor kinase" which is to be a host cell, as described below.

[0051] Examples of the "polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present hystidine kinase" include a polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present hystidine kinase which is derived from a plant-pathogenic filamentous fungus, more specifically, for

example, a polynucleotide having a nucleotide sequence encoding an amino acid sequence selected from the group consisting of:

- (a) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which has an amino acid sequence homology of 95% or more to the amino acid sequence represented by any of SEQ ID NOs: 41, 55 and 68;
- (b) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Fusarium oxysporum-derived cDNA as a template andusing anoligonucleotide having the nucleotide sequence represented by SEQ ID NO: 52 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 53 as primers;
- (c) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Mycospharella tritici-derived cDNA as a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 64 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 65 as primers;
- (d) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Thanapethorus cucumeris-derived cDNA as a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 85 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 86 as primers;
- (e) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is derived from Phytophthora infestans and has the amino acid sequence represented by SEQ ID NO: 90;
- (f) the amino acid sequence represented by SEQ ID NO: 41;
- (g) the amino acid sequence represented by SEQ ID NO: 55; and
- (h) the amino acid sequence represented by SEQ ID NO: 68.
- [0052] One example of a process for producing the transformed cell will be shown below.
  - (1) Preparation of cDNA

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- [0053] First, total RNA is prepared from filamentous fungus, for example, according to the method described in Molecular Cloning 2nd edition authored by J., Sambrook, E., F., Frisch, T., Maniatis. Specifically, for example, a part of a fungal tissue is collected from Neurospora crassa, Botrytinia fuckeliana, Magnaporthe grisea, Phytophthora infestans, Thanatephorus cucumeris, Fusarium oxysporum, Mycospharella tritici, Thanatephorus cucumeris, Thanatephorus cucumeris and the like, the collected tissue is frozen in liquid nitrogen, and is physically ground with a mortar or the like. Then, total RNA may be prepared by the conventional method such as (a) a method of adding a solution containing guanidine hydrochloride and phenol or a solution containing SDS and phenol to the resulting ground material, to obtain total RNA, or (b) a method of adding a solution containing guanidine thiocyanate to the aforementioned ground material, and further adding CsCl, followed by centrifugation, to obtain total RNA. In the procedures, a commercially available kit such as RNeasy Plant Mini Kit (manufactured by QIAGEN) may be also used.
- [0054] Then, the thus prepared total RNA is used to prepare a cDNA. For example, cDNA may be prepared by reacting a reverse transcriptase on the total RNA after an oligo-dT chain or a random primer is annealed to total RNA. In addition, further, a double-stranded cDNA can be prepared, for example, by reacting RNaseH, DNA Polymerase I on said cDNA. In the procedures, a commercially available kit such as SMARTTM PCR cDNA Synthesis Kit (manufactured by Clonech), cDNA Synthesis Kit (manufactured by TAKARA SHUZO Co., Ltd.), cDNA Synthesis Kit (manufactured by Amersham Pharmacia) and ZAP-cDNA Synthesis Kit (manufactured by Stratagene) can be used.
- (2) Cloning
- [0055] When a nucleotide sequence of a desired present histidine kinase is known, a polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kinase can be obtained, for example, from the cDNA prepared as described above, for example, by PCR using as a primer an oligonucleotide having a partial nucleotide sequence of the known nucleotide sequence, or a hybridization method using as a probe an oligonucleotide having a partial nucleotide sequence of the known nucleotide sequence.
- [0056] A polynucleotide having a nucleotide sequence encoding an amino acid sequence of BcOS-1 which is the present histidin kinase can be prepared from a cDNA of Botryotinia fuckeliana, for example, by PCR using as a primer an oligonucleotide having a partial nucleotide sequence of the nucleotide sequence represented by SEQ ID NO: 2, or a hybridization method using as a probe an oligonucleotide having a partial nucleotide sequence of the nucleotide sequence represented by SEQ ID NO: 2.
- [0057] In addition, a polynucleotide having a nucleotide sequence encoding an amino acid sequence of HIK1 which

is the present histidine kinase can be obtained from a cDNA of Magnaporthe grisea, for example, by PCR using as a primer an oligonucleotide having a partial nucleotide sequence of the nucleotide sequence represented by SEQ ID NO: 17, or hybridization method using as a probe an oligonucleotide having a partial nucleotide sequence of the nucleotide sequence represented by SEQ ID NO: 17.

[0058] When a nucleotide sequence of a desired present histidine kinase is unknown, a polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kinase can be obtained by a hybridization method using as a probe an oligonucleotide having a partial nucleotide sequence of the nucleotide sequence of the present histidine kinase, the nucleotide sequence of which is known, or by PCR using as a primer an oligonucleotide designed based on a highly homologous amino acid sequence in plural present histidine kinases, an amino acid sequence of which is known. As the highly homologous amino acid sequence among plural present histidine kinases, amino acid sequences of which are known, for example, there can be exemplified amino acid sequences of a conserved motifs observed in the "repeat sequence region", the "histidine kinase region", the "receiver region" and the like, characterized in the structure of the present histidine kinase.

[0059] More specifically, when the BcOS-1 gene of Botryotinia fuckeliana is obtained by PCR, for example, oligonucleotides designed and synthesized based on nucleotide sequences of about 20bp to about 40bp which are selected from a 5' non-translated region and a 3' non-translated region, respectively, of the nucleotide sequence represented by SEQ ID NO: 2 can be used as a primer set. Examples of the primer set include a set of an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 3 and an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 4. A PCR reaction solution to be used may be prepared by adding a reaction solution designated by a commercially available DNA polymerase or kit as described below to 250ng of a cDNA. The PCR reaction conditions can be appropriately changed depending on a primer set to be used, and examples thereof include the condition of maintaining a temperature at 94°C for 2 minutes, then maintaining a temperature at about 8°C for 3 minutes and, thereafter, repeating around 40 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 30 seconds, then at 55°C for 30 seconds, then at 72°C for 4 minutes, and the condition of repeating 5 to 10 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 5 seconds, then at 72°C for 4 minutes, and further repeating about 20 to 40 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 5 seconds, then at 70°C for 4 minutes. For the procedures, commercially available DNApolymerases contained in Heraculase™ Enhanced DNA Polymerase(manufactured by Toyobo Co.,Ltd.), Advantage cDNA PCR Kit (manufactured by Clonetech), and commercially available kits such as TAKARA Ex Taq (manufactured by TAKARA SHUZO Co., Ltd.), PLATINUMTM PCR SUPERMix (manufactured by Lifetech Oriental), KOD-Plus-(manufactured by Toyobo Co.,Ltd.)and the like can be used.

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[0060] When the hik1 gene of Magnaporthe grisea is obtained by PCR, for example, oligonucleotides designed and synthesized based on nucleotide sequences selected from a 5' non-translation region and a 3' non-translation region, respectively, of the nucleotide sequence represented by SEQ ID NO: 17 can be used as a primer set. Examples of the primer set include a set of an oligonucleotide comprising the nucleotide sequence represented by SEQ ID NO: 18 and an oligonucleotide comprising the nucleotide sequence represented by SEQ ID NO: 19. A PCR reaction solution and the reaction conditions as described above can be used to perform PCR, to obtain the hik1 gene.

[0061] When a gene of the present histidine kinase, a nucleotide sequence of which is not known, is obtained from Fusarium oxysporum, Mycospharella tritici, Thanatephorus cucumeris or Phytophthora infestans, a polynucleotide having a nucleotide sequence encoding a part of an amino acid sequence of the present histidine kinase (hereinafter, referred to as present gene fragment in some cases) can be obtained by the following PCR. As a primer set, for example, a set of oligonucleotides designed and synthesized based on amino acid sequences of a conserved motifs observed in the "repeat sequence region", the "histidine kinase region", the "receiver region" and the like, characterized in the structure of the present histidine kinase, can be used. Examples of the primer set include a primer set of an oligonucleotide having the nucleotide sequence represented by any of SEQ ID NOs: 30 to 34 and an oligonucleotide having the nucleotide sequence represented by any of SEQ ID NOs: 35 to 40.

[0062] Specifically, in the case of Fusarium oxysporum, for example, using an oligonucleotide primer having the nucleotide sequence represented by SEQ ID NO: 33 and an oligonucelotide primer having the nucleotide sequence represented by SEQ ID NO: 38, and using KOD-Plus-(TOYOBO), amplification is performed under the conditions in which a temperature is maintained at 94°C for 2 minutes, and then 35 cycles of incubation are repeated, each cycle comprising maintaining a temperature at 94°C for 15 seconds, then, at 55°C for 30 seconds and, further, at 68°C for 5 minutes. In addition, in the case of Mycospharella tritici, for example, using an oligonucleotide primer having the nucleotide sequence represented by SEQ ID NO: 31 and an oligonucelotide primer having the nucleotide sequence represented by SEQ ID NO: 40, and using KOD-Plus- (TOYOBO), amplification is performed under the conditions in which a temperature is maintained at 94°C for 2 minutes, and 35 cycles of incubation are repeated, each cycle comprising maintaining a temperature at 94°C for 15 seconds, then, at 55°C for 30 seconds and, further, at 68°C for 3 minutes. In addition, in the case of Thanatephorus cucumeris, for example, using an oligonucelotide primer having the nucleotide sequence represented by SEQ ID NO: 30 and an oligonucleotide primer having the nucleotide sequence

represented by SEQ ID NO: 37, and using KOD-Plus-(TOYOBO), amplification is performed under the conditions in which a temperature is maintained at 94°C for 2 minutes, and 35 cycles of incubation are repeated, each cycle comprising maintaining a temperature at 94°C for 15 seconds, then, at 55°C for 30 seconds, further, at 68°C for 1 minute. In addition, in the case of Phytophthora infestans, for example, using an oligonucleotide primer having the nucleotide sequence represented by SEQ ID NO: 31 and an oligonucleotide primer having the nucleotide sequence represented by SEQ ID NO: 37, and using KOD-Plus- (TOYOBO), amplification is performed under the conditions in which a temperature is maintained at 94°C for 2 minutes, and 35 cycles of incubation are repeated, each cycle comprising maintaining a temperature at 94°C for 15 seconds, then, at 55°C for 30 seconds and, further, at 68°C for 1 minute. By such the PCR, a polynucleotide having a nucleotide sequence encoding a part of an amino acid sequence of the present histidine kinase is amplified. A polynucleotide having a nucleotide sequence encoding a full length amino acid sequence of the present histidine kinase can be obtained by RACE method by using, for example, SMART RACE cDNA Amplification Kit (CLONTECH) and primers designed based on a nucleotide sequence of the amplified polynucleotide (present gene fragment).

[0063] When the polynucleotide obtained as described above has revealed a nucleotide sequence encoding a full length amino acid sequence of the present histidine kinase, by PCR using an oligonucelotide having a partial nucleotide sequence of the sequence as a primer, a polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kinase may be also obtained.

[0064] Specifically, when a gene of the present histidine kinase of Fusarium oxysporum (hereinafter, referred to FoOS-1 gene in some cases) is obtained by PCR, for example, oligonucelotides designed and synthesized based on nucleotide sequences selected from 5'-terminal region and a 3'-terminal region, respectively, of the nucleotide sequence represented by SEQ ID NO: 42 can be used as a primer set. Examples of the primer set include a set of an oligonucleotide comprising the nucleotide sequence represented by SEQ ID NO: 52 and an oligonucelotide comprising the nucleotide sequence represented by SEQ ID NO: 53. A PCR reaction solution and the reaction conditions as described above are used to perform PCR, whereby, a polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kinase derived from Fusarium oxysporum can be obtained.

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[0065] In addition, when a gene of the present histidine kinase of Mycospharella tritici (hereinafter, referred to StOS-1 gene in some cases) is obtained by PCR, for example, oligonucleotides designed and synthesized based on nucleotide sequences selected from a 5'-terminal region and a 3'-terminal region, repectively, of the nucleotide sequence represented by SEQ ID NO: 56 can be used as a primer set. Examples of the primer set include a set of an oligonucleotide comprising the nucleotide sequence represented by SEQ ID NO: 64 and an oligonucleotide comprising the nucleotide sequence represented by SEQ ID NO: 65. A PCR reaction solution and the reaction conditions as described above are used to perform PCR, whereby, a polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kinase derived from Mycospharella tritici can be obtained.

[0066] In addition, when a gene of the present histidine kinase of Thanatephorus cucumeris (hereinafter, referred to RsOS-1 gene in some cases) is obtained by PCR, for example, oligonucelotides designed and synthesized based on nucleotide sequences selected from 5'-terminal region and a 3'-terminal region, respectively, of the nucleotide sequence represented by SEQ ID NO: 69 can be used as a primer set. Examples of the primer set include a set of an oligonucelotide comprising the nucleotide sequence represented by SEQ ID NO: 85 and an oligonucleotide comprising the nucleotide sequence represented by SEQ ID NO: 86. A PCR reaction solution and the reaction conditions as described above are used to perform PCR, whereby, a polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kinase derived from Thanatephorus cucumeris can be obtained.

[0067] When a hybridization method is used, cloning can be performed, for example, according to the method described in Molecular Cloning 2nd edition, authored by J., Sambrook, E., F., Frisch, T., Maniatis.

[0068] A probe used to obtain a gene of the present histidine kinase can be obtained by synthesizing a DNA (around about 200 bases to about 500 bases in length) having a partial nucleotide sequence of the nucleotide sequence represented by SEQ ID NO: 2, followed by radioisotope-labeling or fluorescently labeling the DNA according to the conventional method. In such the labeling of a DNA, commercially available kits such as Random Primed DNA Labelling Kit (manufactured by Boehringer), Random Primer DNA Labelling Kit Ver.2 (manufactured by TAKARA SHUZO Co., Ltd.), ECL Direct Nucleic acid Labelling and Detection System (manufactured by Amersham Pharmacia), Megaprime DNA-labelling system (manufactured by Amersham Pharmacia) and the like may be utilized. The thus obtained probe can be used for cloning a gene of the histidine kinase such as the BcOS1-gene of Botrytinia fuckeliana, a nucleotide sequence of which is known, or a gene of the present histidine kinase, a nucleotide sequence of which is unknown.

[0069] Examples of the hybridization condition include the stringent condition, specifically, the condition under which,

in the presence of 6×SSC (0.9 M NaCl, 0. 09 M trisodium citrate), 5×Denhart's solution (0.1% (w/v) Ficoll 400, 0.1% (w/v) polyvinylpyrrolidone, 0.1%BSA), 0.5% (w/v) SDS and 100 µg/ml denatured salmon sperm DNA, or in DIG EASY Hyb solution (Boehringer Manheim) containing 100 µg/ml denatured salmon sperm DNA, a temperature is maintained at 65°C, then a temperature is maintained at room temperature for 15 minutes twice in the presence of 1×SSC (0.15 M NaCl, 0.015 M trisodium citrate) and 0.5%SDS, further, a temperature is maintained at 68°C for 30 minutes in the

presence of 0.1×SSC (0.015 M NaCl, 0.0015M trisodium citrate) and 0.5%SDS.

[0070] Specifically, for example, for obtaining the BcOS-1 gene of Botrytinia fuckeliana, PCR is performed by using a Botrytinia fuckeliana cDNA library phage solution (about 1,000,000 pfu) as a template, and using TAKARA LA taq™ (manufactured by TAKARA SHUZO Co., Ltd.), and using an oligonucleotide comprising the nucleotide sequence represented by SEQ ID NO: 9 and an oligonucleotide comprising a nucleotide sequence complementary to the nucleotide sequence represented by SEQ ID NO: 10 as a primer set, whereby, a DNA for a probe is amplified, which may be collected. A PCR reaction solution to be used may be prepared by adding a reaction solution designated by a kit as described above to 250ng of a DNA library. Examples of the PCR reaction condition include the condition under which amplification is performed by maintaining a temperature at 94°C for 2 minutes, then at 8°C for 3 minutes, and repeating 40 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 30 seconds, then, at 55°C for 30 seconds and, then, at 68°C for 5 minutes. Then, a probe labeled with 32P can be prepared by using the amplified and obtained DNA as a template, and using Megaprime DNA-labelling system (Amersham Pharmacia) and using a reaction solution designated by the kit. The thus prepared probe is used to perform colony hybridization according to the conventional method, in which a temperature is maintained at 65°C in the presence of 6×SSC (0.9M NaCl, 0.09M trisodium citrate, 5×Denharp's solution (0.1%(w/v) Ficoll 400, 0.1% (w/v) polyvinylpyrrolidone, 0.1%BSA), 0.5%(w/v) SDS and 100 µg/ml denatured Salmon sperm DNA, or in DIG EASY Hyb solution (Boehringer Mannheim), containing 100 µg/ml denated Salmon sperm DNA, then, a temperature is maintained at room temperature for 15 minutes twice in the presence of 1×SSC (0.15 M NaCl, 0.015M trisodium citrate) and 0.5%SDS and, further, a temperature is maintained at 68°C for 30 minutes in the presence of 0.1×SSC (0.015 M NaCl, 0.0015 M sodium citrate) and 0.5%SDS, whereby, a clone which hybridizes with the probe can be obtained.

[0071] In addition, a gene of the present histidine kinase having a known nucleotide sequence may be also prepared by performing chemical synthesis of a nucleic acid, for example, according to the conventional method such as a phosphite triester method (Hunkapiller, M. et al. Nature 310, 105, 1984), based on the known nucleotide sequence. [0072] The thus obtained polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kinase may be cloned into a vector according to the conventional method described in "Molecular Cloning: A Laboratory Manual 2nd edition" (1989), Cold Spring Harbor Laboratory Press, "Current Protocols In Molecular Biology" (1987), John Wiley & Sons, Inc. ISBNO-471-50338-X or the like. Examples of the vector to be used include pBlueScript II vector (manufacturedbyStratagene), pUC18/19 vector (manufactured by TAKARA SHUZO Co., Ltd.), TA Cloning vector (manufactured by Invitrogen) and the like.

[0073] A nucleotide sequence of the cloned gene may be confirmed by the Maxam Gilbert method (described in Maxam, A.M. &W.Gilbert, Proc. Natl. Acad. Sci. USA, 74, 560, 1977 etc.) or the Sanger method (described in Sanger, F. & A. R. Coulson, J. Mol. Biol., 94, 441, 1975, Sanger, F, & Nicklen and A.R.Coulson., Proc. Natl. Acad. Sci. USA, 74, 5463, 1977 etc.). For the procedures, commercially available kits such as Termo Seqenase II dye terminator cycle sequencing kit (manufactured by Amersham Pharmacia), Dye Terminator Cycle Sequencing FS Ready Reaction Kit (manufactured by PE Biosystems Japan) and the like can be used.

## (3) Construction of expression vector

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[0074] An expression vector of a polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kinase may be constructed by a conventional method (for example, method described in J. Sambrook, E., F., Frisch, T., Maniatis, Molecular Cloning 2nd edition, published by Cold Spring Harbor Laboratory Press etc.).

[0075] For example, Apolynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kinase may be incorporated into a vector which can be utilized in a host cell to be transformed, for example, a vector which contains genetic information required to be replicable in a host cell, can replicates autonomously, can be isolated and purified from a host cell, and has a detectable marker (hereinafter referred to as basic vector in some cases). As the basic vector, specifically, when a bacterium such as Escherichia coli is used as a host cell, for the example, a plasmid pUC119 (manufactured by TAKARA SHUZO Co., Ltd.), phagemid pBluescriptII (manufactured by Stratagene) and the like may be used. When yeast is used as a host cell, for example, plasmids pACT2 (manufactured by Clontech), p415 CYC (ATCC87382), p415 ADH (ATCC87374) and the like may be used. When a plant cell is used as a host cell, for the example, a plasmid pBl221 (Clontech) and the like may be used.

[0076] An expression vector which can express a polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kinase in a host cell can be constructed by incorporating into a basic vector a polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kinase upstream of which a promoter functional in a host cell is operably linked. Herein, the "operably linked" means that the promoter and a polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kanase are ligated so that the polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kinase is expressed under control of the promoter in a host cell. Examples of a promoter functional

in a host cell include, when a host cell is Escherichia coli, a promoter of a lactose operon (lacP) a promoter of tryptphan operon (trpP), a promoter of an arginine operon (argP), a promoter of a galactose operon (galP), tac promoter, T7 promoter, T3 promoter of Escherichia coli, a promoter of  $\lambda$  phage ( $\lambda$ -pL,  $\lambda$ -pR) and the like. In addition, when a host cell is yeast, examples include an ADH1 promoter, a CYC1 promoter and the like. The ADH1 promoter can be prepared, for example, by the conventional genetic engineering method from a yeast expression vector p415 ADH (ATCC87374) harboring an ADH1 promoter and a CYC1 terminator. The CYC1 promoter can be prepared by the conventional genetic engineering method from p415CYC (ATCC87382). Examples of the promoter include, when a host cell is a plant cell, a nopaline synthase gene (NOS) promoter, an octopinesynthasegene (OCT) promoter, acauliflowermosaicvirus (CaMV)-derived 19S promoter, a CaMV-derived 35S promoter and the like.

[0077] In addition, when a polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kinase is incorporated into a vector already harboring a promoter functional in a host cell, a gene of the present histidine kinase may be inserted into downstream of the promoter so that a promoter harbored by the vector and a gene of the present histidine kinase are operably linked. For the example, the aforementioned yeast plasmid p415 ADH has an ADH1 promoter and, when a gene of the present histidine kinase is inserted downstream of an ADH1 promoter of the plasmid, an expression vector which can express a gene of the present histidine kinase in a budding yeast such as Saccharomyces cerevisiae AH22 (IFO10144 and TM182 (Maeda, T. et al. (1994) Nature 369:242-245) can be constructed.

### (4) Preparation of transformed cell

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[0078] By introducing the constructed expression vector into a host cell according to the conventional method, a transformed cell expressing the present histidine kinase can be prepared. As a host cell used for preparing such the transformed cell, for example, there are bacterium, yeast, plant cell and the like. As the bacterium, for example, there are Escherichia coli, Vibrio harveiy and the like. As the yeast, there are budding yeast and diving yeast. More specifically, for example, there are yeasts belonging to genus Saccharomyces, genus Shizosaccharomycess the like. As a plant cell, for example, there is a plant cell such as Arabidopsis thaliana and the like.

[0079] As a method of introducing an expression vector into the aforementioned host cell, the conventional introducing method can be applied depending on a host cell to be transformed. For example, when bacterium is used as a host cell, the expression vector can be introduced into a host cell by the conventional introducing method such as a calcium chloride method and an electroporation method described in Moleculer Cloning (J. Sambrook et al., Cold spring Harbor, 1989). When yeast is used as a host cell, for example, the expression vector can be introduced into a host cell using Yeast transformation kit (Clontech) based on a lithium method. In addition, when a plant cell is used as a host cell, for example, the expression vector can be introduced into a host cell using the conventional introducing method such as an Agrobacterium infection method (JP-B No.2-58917 and JP-A No.60-70080), an electroporation method into a propoplast (JP-ANo. 60-251887 and JP-ANo. 5-68575) and a particle gun method (JP-A No.5-508316 and JP-A No.63-258525).

(Intracellular signal transduction system regarding present histidine kinase)

[0080] In the present invention, in order to measure an amount of intracellular signal transduction from the present histidine kinase expressed in the transformed cell prepared as described above or an index value having the correlation therewith, an intracellular signal transduction system originally contained in a host cell used for preparing the transformed cell may be utilized. Examples of the intracellular signal transduction system which can be utilized include an intracellular signal transduction system regarding osmolarity responses of the aforementioned budding yeast, an intracellular signal transduction system regarding cell cycle progression and oxidative stress response of fission yeast, an intracellular signal transduction system regarding control of expression of capsular polysaccharide biosynthesis operon in Escherichia coli, an intracellular signal transduction system regarding control of cell density-sensitive luminescence of bioluminescent marine microorganism Vibrio harveyi, an intracellular signal transduction system regarding cytokinin response of Arabidopsis thaliana and the like.

[0081] When the aforementioned expression vector of the present histidine kinase is introduced using the "cell deficient in at least one hybrid-sensor kinase" as a host cell used for preparing such the transformed cell, the produced present histidine kinase functions in place of deleted hybrid-sensor kinase, and intracellular signal is transmitted. In the case where a test substance is contacted with the transformed cell, when signal transduction from the present histidine kinase is inhibited by the test substance, change in an amount of growth of the transformed cell, change in morphology of the transformed cell, change in a shape of the transformed cell, change in an amount of biosynthesis of a particular substance in the cell, change in an amount of metabolism of a particular substance in the cell and the like occur in some cases. In such the cases, an antifungal activity of the test substance acting on the present histidine kinase can be measured using change in an amount of growth of the transformed cell, change in morphology, change

in shape, change in an amount of biosynthesis of a particular substance in a cell, change in an amount of metabolism of a particular substance and the like as an index.

[0082] On the other hand, when at least one intrinsic hybrid-sensor kinase is not deleted in a host cell used for preparing a transformed cell, there are both of signal transduction from intrinsic hybrid-sensor kinases and intracellular signal transduction from the introduced present histidine kinase in intracellular signal transduction of the transformed cell. Change in an amount of growth of the transformed cell, change in morphology, change in shape, change in am amount of biosynthesis of a particular substance in the cell, change in an amount in metabolism of a particular substance in the cell and the like reflecting an amount of intracellular signal transduction from the introduced present histidine kinase become smaller by the influence of an amount of intracellular signal transduction from intrinsic hybrid-sensor kinase. In the present invention, by using a host cell deficient in at least one intrinsic hybrid-sensor kinase, since change in an amount of growth of the transformed cell, change in morphology, change in shape, change in an amount of biosynthesis of a particular substance in the cell, change in an amount of metabolism of particular substance in the cell and the like reflecting an amount of intracellular signal transduction from the introduced present histidine kinase become larger, the sensitivity of the transformed cell to an antifungal compound is enhanced. Like this, the transformed cell with the enhanced sensitivity to an antifungal compound is useful for assaying the antifungal activity of a test substance and searching an antifungal compound using the assay.

[0083] Specifically, when the present histidine kinase is introduced in a Saccharomyces cerevisiae strain deficient in hybrid-sensor kinase SLN1 (Maeda, T. et al. Nature:369 242-245 (1994)), the present histidine kinase performs signal transduction in place of deficient SLNI, whereby, an amount of intracellular signal transduction from the introduced present histidine kinase can be detected more clearly using, an amount of growth of host cell as an index. That is, when the test substance acts on the present histidine kinase, and an amount of signal transduction from the present histidine kinase in a host cell is changed, it can be clearly measured as change in an amount of growth of the trans formed budding yeast. In addition, an Escherichia coli strain deficient in a hybrid-sensor kinase RcsC, a fission yeast strain deficient in PHK1 to PHK3 involved in control of cell cycle progression, a Vibrio harveyi strain deficient in LuxN associated with control of cell density-sensitive luminescence and an Arabidopsis thaliana strain deficient in cytokinin receptor CRE1 can be exemplified as one preferable aspect of the "cell deficient in at least one hybrid-sensor kinase".

(Method of assaying antifungal activity of test substance)

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[0084] In a method of assaying the antifungal activity of a test substance, an embodiment of a first step of culturing a transformed cell in which a polynucleotide having a nucleotide sequence encoding an amino acid sequence of the present histidine kinase introduced in the presence of a test substance includes a method of contacting a test substance with the transformed cell by culturing the transformed cell in a medium containing the test substance. Culturing the transformed cell may be any form of liquid culturing in which the cell is cultured in a liquid medium, solid culturing in which the cell is cultured on a solid medium prepared by adding agar or the like to liquid medium, and the like. The concentration of a test substance in the medium is, for example, about 1 nin to about 1 mM, preferably about 10 nm to about 100 µM. A culturing time is, for example, about 1 hour or longer and around 3 days, preferably about 25 hours to around 2 days. When the antifungal activity of a test substance is assayed, as a medium containing a test substance, an antifungal compound-free medium may be used.

[0085] An amount of intracellular signal transduction from the present histidine kinase expressed in a transformed cell cultured in the first step or an index value having the correlation therewith is measured. And, the antifungal activity of a test substance is assayed based on a difference between an amount of intracellular signal transduction or an index value having the correlation therewith measured in the second step and a control. For example, the antifungal activity of the test substance can be assessed based on a difference obtained by comparing amounts of intracellular signal transduction or index values having the correlation therewith, which are measured as described above in sections in which different two or more substances (for example, it is preferable that among different two or more substances, at least one substance has no antifungal activity) are independently used, respectively, as a test substance.

[0086] Specifically, for example, when a transformed cell prepared by using, as a host cell, the TMI82 (SLNI\(\Delta\)) strain (Maeda T. et al. Nature:369 242-245 (1994)) which is a SLNI gene-deficient strain in which the PTP2 Tyrosine phosphatase gene (Ota et al, Proc.N.A.sic.USA, 89, 2355-2359 (1992)) introduced (that is, a transformed cell having the function that cell growth is directly controlled by transduction of an intracellular signal from the present histidine kinase) is used, the antifungal activity can be measured by using, as an index, an amount of growth of the transformed cell in a medium (agar medium or liquid medium) using glucose as a carbon source, for example, Glu-Ura-Leu medium. When a medium in which a test substance is added to the Glu-Ura-Leu medium (medium containing no antifungal compound) is used, a test substance inhibiting growth of the transformed cell can be assessed to have the antifungal activity. In addition, as a control, it is enough to examine that growth of the transformed cell in a medium using galactose in place of glucose as a carbon source, for example, Gal-Ura-Leu medium is observed regardless of the presence or the absence of test substance.

[0087] When a transformed cell prepared by using, as a host cell, fission yeast which is PHK1, PHK2 and PHK3 gene-deficient strain (that is, a transformed cell in which cell cycle progression is directly regulated by transduction of an intracellular signal from the histidine kinase) is used, cell division of the fission yeast may be observed under a microscope. When a medium in which a test substance is added to a medium containing no substance having the antifungal activity is used, a test substance which shortens a cell length of a dividing cell of the transformed cell can be assessed to have the antifungal activity.

[0088] When a transformed cell prepared by using, as a host cell, RcsC gene-deficient Escherichia coli in which cps-LacZ introduced is used, color development of X-Gal may be observed in an agar medium or a liquid medium (Suzuki et al. Plant Cell Physiol. 42:107-113(2001)). When a medium in which a test substance is added to a medium containing no substance having the antifungal activity is used, a test substance which can make the transformed cell develop blue can be assessed to have the antifungal activity.

[0089] In addition, when a transformed cell prepared by using, as a host cell, LuxN gene-deficient V, harveyi (i.e. a transformed cell in which bioluminescence is directly regulated by transduction of an intracellular signal from the present histidine kinase) is used, the fluorescent light emitted by the transformed microorganism may be observed. When a medium containing a test substance and not containing a substance having the antifungal activity is used, a test substance which make the transformed cell possible to emit the fluorescent light can be assessed to have the antifungal activity.

[0090] Further, a substance having the antifungal activity can be also searched by selecting an antifungal compound based on the antifungal activity assessed by the aforementioned assaying method.

Effects of the invention

[0091] The present invention can provide a transformed cell with the enhanced sensitivity to an antifungal compound, a method of assaying the antifungal activity of a test substance using the transformed cell, and a method of searching an antifungal compound using the method.

Examples

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[0092] The present invention is further described in the following Examples, which are not intended to restrict the invention.

Example 1

Isolation of Botryotinia fuckeliana BcOS-1 gene

-[0093] Total RNA was prepared from Botryotinia fuckeliana. 100 mg of a hypha of Botryotinia fuckeliana strain Bc-16 grown on a potato dextrose agar medium (PDA medium manufactured by NISSUI Pharmaceutical Co., Ltd.) was scratched off, and this was ground in liquid nitrogen using a mortar and a pestle. A RNA was prepared from frozen ground powder using RNeasy Plant Mini Kit (QIAGEN). A frozen ground powder together with liquid nitrogen was transferred to a 50 ml sample tube and, after liquid nitrogen was all volatilized, a solution obtained by adding 10 μL of mercaptoethanol per 1 ml of a buffer RLC attached to kit was added, followed by stirring. Further, ground powder was well dispersed by a few of pipettings, and was incubated at 56°C for 3 minutes. Thereafter, the solution containing ground powder was supplied to QIAshredder spin column attached to the kit, and centrifuged at  $8,000 \times g$  for 2 minutes. The filtration supernatant was transferred to a fresh sample tube, a 0.5-fold volume of 99.5% ethanol was added thereto, and the material was well mixed by pipetting. This mixture was supplied to RNeasy mini spin column attached to the kit, and centrifuged at 8, 000×g for 1 minute. The filtrate was discarded, the residue was added 700 μL of a buffer RWI attached to the kit, and centrifuged at 8,000 imes g for 1 minute, and the filtrate was discarded. Further, the residue was added 500  $\mu$ L of a buffer RPE attached to the kit, centrifuged at 8,000imesg for 1 minute, and the filtrate was discarded. This procedure was repeated twice. Finally, an upper filter part was transferred to a fresh sample tube, supplied 30 µL of RNase-free sterilized water attached to the kit, and centrifuged at 8,000×g for 1 minute, and total RNA was dissolved out into the filtrate. This dissolution procedure was repeated twice. The concentration of the resulting total RNA solution was obtained from the absorbance at 260 nm to be 322 µg/ml.

[0094] Then, a cDNA was synthesized using ThermoScript RT-PCR System (Invitrogen) while employing total RNA as a template. A solution in which 2.7  $\mu$ L of total RNA and 6.3  $\mu$ L of sterilized distilledwater were mixed into 1.0  $\mu$ L of 50 mM Oligo (dt)  $_{20}$  attached to the kit and 2.0  $\mu$ L of 10 mM dNTP Mix was treated at 65°C for 5 minute, and then rapidly cooled on ice. To this solution were added 4  $\mu$ L of 5×cDNA Synthesis Buffer attached to the kit, 1  $\mu$ L of 0.1M DTT, 1  $\mu$ L of RNase OUT, 1  $\mu$ L of ThermoScript RT and 1  $\mu$ L of sterilized distilled water, to react them at 50°C for 60 minutes and, thereafter, the reaction was stopped by heating treatment at 85°C for 5 minutes. Further, a RNA of a

template was degraded by adding 1  $\mu$ L of RNaseH attached to the kit to this reaction solution and maintained a temperature at 37°C for 20 minutes, to obtain a cDNA.

[0095] A DNA having a nucleotide sequence encoding an amino acid sequence of Botryotinia fuckiliana BcOS-1 (hereinafter, referred to as BcOS-1 DNA in some cases) was amplified by PCR using this cDNA as a template. Using an oligonucleotide comprising the nucleotide sequence represented by SEQ ID NO: 3 and an oligonucelotide consisting of the nucleotide sequence represented by SEQ ID NO: 4 as a primer, a PCR was performed to amplify a DNA having the nucleotide sequence represented by SEQ ID NO: 2. The PCR was performed using KOD-Plus-(TOYOBO) under the amplifying conditions that a temperature wasmaintained at 94°C for 2 minutes and, thereafter, 35 cycles of incubation were repeated, each cycle comprising maintaining a temperature at 94°C for 15 seconds, then, at 55°C for 30 seconds, then, at 68°C for 6 minutes. ThePCRreactionsolution (50  $\mu$ L) was prepared by adding 2  $\mu$ L of the aforementioned cDNA, 5  $\mu$ L of 10×Buffer, 5  $\mu$ L of 2 mM dNTPs, 2  $\mu$ L of 25 mM MgSO<sub>4</sub>, each 1  $\mu$ L of 10  $\mu$ M oligonucleotide primers, 33  $\mu$ L of sterilized distilled water and 1  $\mu$ L of KOD-Plus-. After the reaction, a part of the reaction solution was separated by 0.8% agarose gel electrophoresis, and stained with ethidium bromide. It was confirmed that about 4 kb of a DNA (BcOS-1 DNA) was amplified.

#### Example 2

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Construction of expression plasmid of Botryotinia fuckeliana BcOS-1 gene and preparation of transformed budding yeast

[0096] BcOS-1 DNA was cloned into a shuttle vector p415ADH (ATCC87312) replicable in yeast and Escherichia coli. About 4 kb of the aforementioned DNA (BcOS-1 DNA) was purified from the PCR reaction solution prepared in Example 1 using QIAquick PCR Purification Kit (QIAGEN) according to the attached manual. About 4 kb of the purified DNA (BcOS-1 DNA) was digested with restriction enzymes Spel and Pstl and, on the other hand, the shuttle vector p415ADH was also digested with restriction enzymes Spel and Pstl and, thereafter, each of which was separated by 0. 8% agarose gel electrophoresis, and apart of the gel containing a desired DNA was excised. The BcOS-1 DNA digested with Spel and Pstl and the shuttle vector digested with Spel and Pstl were recovered from the gel using QIAquick Gel Extraction Kit (QIAGEN) according to the attached manual. The aforementioned BcOS-1 DNA was inserted between Spel site and Pstl site in the multicloning site of the shuttle vector using Ligation Kit Ver. 2 (TaKaRa) according to the attached manual, to construct an expression plasmid pADHBcOS1. A nucleotide sequence of the resulting expression plasmid was analyzed with a DNA sequencer (Model 3100, Applied Biosystems) after a sequencing reaction using BigDye terminator v3.0 Cycle Sequencing Ready Reaction Kit (Applied Biosystems) according to the attached manual. The sequencing reaction was performed using an oligonucleotide consisting of the nucleotide sequence represented by any of SEQ ID NOs: 5 to 12 as a primer under the amplifying conditions that 30 cycles were repeated, each cycle comprising maintaining a temperature at 96°C for 10 seconds, then, at 50°C for 5 seconds, then, -at 60°C for 4 minutes. As a result, the nucleotide sequence represented by SEQ ID NO: 2 was obtained, and it was confirmed that the expression plasmid pADHBcOS1 harbored a DNA having a nucleotide sequence encoding an amino acid sequence of BcOS-1.

[0097] The prepared expression plasmid pADHBcOS1 was introduced into each of budding yeast (Saccharomyces cerevisiae) AH22 strain (IFO10144) and TM182 strain (Maeda T. et al. (1994) Nature vol. 369, pp242-245) according to the method described in Geitz RD & Woods RA (1994) Molecular Genetics of Yeast: Practical Approaches ed. Johnson JA, Oxford University Press pp124-134. By utilizing disappearance of leucine auxotrophy in the resulting transformed budding yeast, the transformed budding yeast AH 22 strain (AH22-BcOS1) was selected on a Glu-Leu agar medium, and the transformed budding yeast TM182 strain (TM182-BcOS1) was selected on a Gal-Ura-Leu agar medium. It was confirmed that the resulting TM182-BcOS1 grows even when transplanted to a Glu-Ura-Leu medium.

#### Example 3

Antifungal compound sensitivity test of transformed budding yeast TM182-BcOS1

[0098] The transformed budding yeast AH22-BcOS1 prepared in Example 2 was cultured while shaking at 30°C for 18 hours in a Glu-Leu medium. As a control, the AH22 strain was similarly cultured while shaking at 30°C for 18 hours in a Glu medium. The absorbance at 600 nm of each of the grown transformed budding yeasts in a cell suspension was measured, and a cell suspension diluted with sterilized distilled water to the absorbance of 0.1 was prepared. Further, a cell suspension in which the aforementioned cell suspension of the transformed budding yeast AH22-BcoS1 was diluted 200-fold with a Glu-Leu medium, and a cell suspension in which the aforementioned cell suspension of the AH22 strain was diluted 200-fold with a Glu medium were prepared. A solution in which each of Compounds (1) to (3) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 60 ppm, a solution in which each of Compounds

(4) and (5) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 2000 ppm, and a solution in which each of Compounds (6) and (7) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 20 ppm were prepared. and two microplates were prepared in which each 2.0 µL per well of the Compound solution and DMSO as a control were dispensed into two wells. In one microplate among them, each 200 µL of cell suspensions of the transformed budding yeast AH22-BcOS1 which had been prepared by dilution as described above was dispensed, and cultured by allowing to stand at 30°C for 48 hours. In another microplate, each 200 µL of the cell suspensions of the control yeast AH22 strain which had been prepared by dilution as described above was dispensed, and cultured by allowing to stand at 30°C for 48 hours. After culturing, the absorbance at 600 nm of each well was measured with a microplate reader. [0099] Similarly, the transformed budding yeast TM182-BcOS1 prepared in Example 2 was cultured at 30°C for 18 hours in a Glu-Ura-Leu medium. The absorbance at 600 nm of the grown transformed budding yeast in a cell suspension was measured, and a cell suspension diluted with sterilized distilled water to the absorbance of 0.1 was prepared. Further, a cell suspension in which the aforementioned cell suspension of the transformed budding yeast TM182-BcOS1 was diluted 200-foldwith a Glu-Ura-Leu medium and, as a control, a cell suspension in which the aforementioned cell suspension was diluted 200-fold with a Gal-Ura-Leu medium were prepared. A suspension in which each of Compounds (1) to (3) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 60 ppm, a solution in which each of Compounds (4) and (5) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 2000 ppm, and a solution in which each of Compounds (6) and (7) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 20 ppm were prepared, and two microplates were prepared in which each 2.0 µL per well of the Compound solution and DMSO as a control were dispensed into two wells. In one microplate among them, each 200 µL of cell suspensions of the transformed budding yeast TM182-BcOSI which had been prepared by dilution with a Glu-Ura-Leu medium as described above was dispensed, and cultured by allowing to stand at 30°C for 67 hours. In another microplate, as described above, as a control, each 200 µL of the cell suspensions of the transformed budding yeast TM182-BcOSI which had been prepared by dilution with a Gal-Ura-Leumedium was dispensed, and cultured by allowing to stand at 30°C for 67 hours. After culturing, the absorbance at 600 nm of each well was measured with a microplate reader.

[0100] Degree of growths of both of the transformed budding yeasts cultured under the presence of each of Compound (1) to (7) and budding yeast as a control therefor are shown in Table 1. Degree of growths of both of the transformed budding yeasts and budding yeasts as a control therefor are expressed by a relative value in percentage, letting the absorbance at 600 nm in a well having the concentration of the aforementioned Compound of 0 ppm to be 100. It was confirmed that an inhibiting degree of growth of TM182-BcOSI by each test substance was grater than an inhibiting degree of growth of AH22-BcOSI by each test substance, and the TM182-BcOS1 was a transformed cell with the enhanced sensitivity to an antifungal compound as compared with AH22-BcOS1.

Table 1

Test substance (final concentration)	Degree of growth of budding yeast (%)			
	AH22	AH22-Bc TM182-BcOS1  n Glu-Leu medium Gal-Ura-Leu medium		
	Glu medium		Gal-Ura-Leu medium	Gal-Ura-Leu medium
Compound (1) (0.6ppm)	99	90	99	9
Compound (2) (0.6 ppm)	99	92	98	11
Compound (3) (0.6ppm)	98	93	98	10
Compound (4) (20 ppm)	96	45	102	10
Compound (5) (20 ppm)	97	79	103	48
Compound (6) (0.2 ppm)	99	81	99	8
Compound (7) (0.2 ppm)	101	94	99	11

Example 4

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Isolation of Botryotinia fuckeliana mutant BcOS-1 gene exhibiting resistance to dicarboxyimide antifungal compound

[0101] A DNA having a nucleotide sequence encoding an amino acid sequence of Botryotinia fuckeliana mutant BcOS-1 (Oshima, M. et al. (2002) Phytopathology 92, pp75-80) exhibiting resistance to a dicarboxyimide antifungal compound (hereinafter, referred to as mutant BoOS1 DNA in some cases) was prepared by PCR using the cDNA prepared in Example 1 as a template. A first time PCR was performed using, as a primer, an oligonucelotide consisting

of the nucleotide sequence represented by SEQ ID NO: 15 and an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 4, and a DNA having a nucleotide sequence represented by base numbers 1081 to 3948 of the nucleotide sequence represented by SEQ ID NO: 14 was amplified. The PCR was performed using KOD-Plus-(TOYOBO) under the amplifying conditions that a temperature was maintained at 94°C for 2 minutes and, thereafter, 35 cycles were repeated, each cycle comprising maintaining a temperature at 94°C for 15 seconds, then, at 55°C for 30 seconds, then, at 68°C for 6 minutes. The PCR reaction solution (50  $\mu$ L) was prepared by adding 2  $\mu$ L of the aforementioned cDNA, 5  $\mu$ L of 10×Buffer, 5  $\mu$ L of 2 mM dNTPs, 2  $\mu$ L of 25 mM MgSO<sub>4</sub>, each 1  $\mu$ L of 10  $\mu$ M oligonucleotide primers, 33  $\mu$ L of sterilized distilled water and 1  $\mu$ L of KOD-Plus-. After the reaction, a second PCR was performed using an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 3 and 1  $\mu$ L of the first time PCR reaction solution while using the cDNA prepared in Example 1 as a template. The reaction conditions were the same as those of the first time PCR and after the reaction, a part of the reaction solution was separated by 0.8% agarose gel electrophoresis, and stained with ethidium bromide. It was confirmed that about 4 kb of the DNA (mutant BcOS-1 DNA) was amplified.

#### 15 Example 5

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Construction of expression plasmid of Botryotinia fuckeliana BcOS-1 mutant gene exhibiting resistance to dicarboxyimde antifungal compound and preparation of transformed budding yeast

[0102] First, the mutant BcOS-1 DNA was cloned into a vector pBluescript II SK(+) (TOYOBO). About 4 kb of the DNA (mutant BcOS-DNA) was purified from the second time PCR reaction solution prepared in Example 4 using QIAquick PCR Purification Kit (QIAGEN) according to the attached manual. About 4 kb of the purified DNA (mutant BcOS-1 DNA) was digested with restriction enzymes Spel and Pstl and, on the other hand, the vector pBluescript II SK(+) was also digested with restriction enzymes Spel and Pstl, each of which was separated by 0.8% agarose gel electrophoresis, and a part of the gel containing a desired DNA was excised. The mutant BcOS-1 DNA digested with Spel and Pstl and the vector pBluescript II SK(+) digested with Spel and Pstl were recovered from the gel using QIAquick Gel Extraction Kit (QIAGEN) according to the attached manual. The aforementioned mutant BcOS-1 DNA was inserted between Spel site and Pstl site in the multicloning site of the vector pBluescript II SK (+) using Ligation Kit Ver.2 (TaKaRa) according to the attached manual, to construct a plasmid pBcOS1-I 365S. A nucleotide sequence of the resulting plasmid was analyzed with a DNA sequencer (Model 3100, Applied Biosystems) after a sequencing reaction using BigDye terminator v3.0 Cycle Sequence FS Ready Reaction Kit (Applied Biosystems) according to the attached manual. The sequencing reaction was performed by using an oligonucleotide consisting of the nucleotide sequences represented by any of SEQ ID NOs: 7 to 12 as a primer under the amplifying conditions that 30 cycles were repeated, each cycle comprising maintaining a temperature at 96°C for 10 seconds, then, at 50°C for 5 seconds, then, at 60°C for 4 minutes. As a result, the nucleotide sequence represented by SEQ ID NO: 14 was obtained and it was confirmed that the plasmid pBcOS1-I 365S harbored the mutant HcOS-1 DNA.

[0103] The mutant BcOS-1 DNA contained in the thus prepared plasmid pBcOS1-I365S was cloned into a shuttle vector p415ADH replicable in yeast and Escherichia coli, to construct an expression plasmid. The plasmid pBcOS1-I365S was digested with restriction enzymes Spel and Pstl and, on the other hand, the shuttle vector p415ADH was also digested with restriction enzymes Spel and Pstl. These were separated by 0.8% agarose gel electrophoresis, respectively, each of gel parts containing the mutant BcOS-1 DNA digested with Spel and Pstl and the shuttle vector p415ADH digested with Spel and Pstl was excised, and the mutant BcOS-1 DNA and the shuttle vector were recovered from the gel using QIAquickGel Extraction Kit (QIAGEN) according to the attached manual. The mutant BcOS-1 DNA was inserted between Spel site and Pstl site in the multicloning site of the shuttle vector using Ligation Kit Ver.2 (TaKa-Ra) according to the attached manual, to construct an expression plasmid pADHBcOS1-I365S. A nucleotide sequence of the resulting expression plasmid was analyzed with a DNA sequencer (Model. 3100, Applied Biosystems) after a sequencing reaction using BigDye terminator v3.0 Cycle Sequence FS Ready Reaction Kit (Applied Biosystems) according to the attached manual. The sequencing reaction was performed by using an oligonucleotide consisting of the nucleotide sequence represented by any of SEQ ID NOs: 5 to 12 as a primer under the amplifying conditions that 30 cycles of incubation were repeated, each cycle comprising maintaining a temperature at 96°C for 10 seconds, then, at 50°C for 5 seconds, then, at 60°C for 4 minutes. As a result, the nucleotide sequence represented by SEQ ID NO: 14 was obtained, and it was confirmed that the expression plasmid pADHBcOS1-I365S harbored a DNA having a nucleotide sequence encoding an amino acid sequence of the mutant BcOS-1.

[0104] The prepared expression plasmid pADHBcOS1-I 365S was introduced into the budding yeast TM182 strain according to the method described in Example 2. By utilizing disappearance of leucine auxotrophy in the resulting transformed budding yeast, the transformed budding yeast TM182 strain (TM182-BcOS1-I365s) was selected on a Gal-Ura-Leu agarose medium. It was confirmed that the resulting TM182-BcOS1-I365S grows even when transplanted to a Glu-Ura-Leu medium.

### Example 6

Antifungal compound sensitivity test of transformed budding yeast TM182-BcOS1-I-365S

[0105] The transformed budding yeast TM182-BcOS1-I365S prepared in Example 5 was cultured at 30°C for 18 hours in a Glu-Ura-Leu medium. The absorbance at 600 nm of a cell suspension of the grown transformed budding yeast was measured, and a cell suspension diluted with sterilized distilled water to the absorbance of 0.1 was prepared. Further, a cell suspension in which the aforementioned cell suspension of the transformed budding yeast TM182-BcOS1-I 365S was diluted 200-fold with a Glu-Ura-Leu medium and, as a control, a cell suspension in which the cell suspension was diluted 200-fold with a Gal-Ura-Leu medium were prepared. A solution in which each of Compound (1) to (3) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 60 ppm, a solution in which each of Compounds (4) and (5) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 2000 ppm, and a solution in which each of Compounds (6) and (7) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 20 ppmwere prepared, and two microplates were prepared in which each 2.0 µL per well of the Compound solution and DMSO as a control were dispensed into two wells. In one microplate among them, each 200 μL of cell suspensions of the transformed budding yeast TM182-BcOS1-I365S which had been prepared by dilution with a Glu-Ura-Leumedium as described above was dispensed, and cultured by allowing to stand at 30°C for 67 hours. In another microplate, as a control, each 200 µL of cell suspensions of the transformed budding yeast TM182-BcOS1-I 365S which had been prepared by dilution with a Gal-Ura-Leu medium was dispensed, and cultured by allowing to stand at 30°C for 67 hours. After culturing, the absorbance at 600 nm of each well was measured with a microplate reader.

[0106] Degrees of growths of both of the transformed budding yeasts cultured under the presence of Compounds (1) to (7) and budding yeast as a control therefor are shown in Table 2. Degrees of growths of both of the transformed budding yeasts and budding yeast as a control are expressedby a relative value inpercentage, letting the absorbance at 600 nm at the concentration of the Compound of 0 ppm to be 100. It was confirmed that an inhibiting degree of growth of the transformed budding yeast TM182-BcOS1-I 3655 by each test substance was grater than an inhibiting degree of growth of the transformed budding yeast AH22-BcOS1-I 3655 by each test substance, and the transformed budding yeast TM182-BcOS1-I 365S was a transformed cell with the enhanced sensitivity to an antifungal compound as compared with the transformed budding yeast AH22-BcOS1-I365S.

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Table 2

	Degree of growth of budding yeast (%)				
	AH22	AH22-Bc OS1-I36 5S TM182-BcOS1-I365	TM182-BcOS1-I365S	S	
Test substance (final concentration)	Glu medium	Glu-Leu medium	Gal-Ura-Leu medium	Glu-Ura-Leu medium	
Compound (1) (6 ppm)	88	68	99	9	
Compound (2) (6 ppm)	91	81	88	11	
Compound (3) (6 ppm)	87	75	92	9	
Compound (4) (20 ppm)	96	83	101	41	
Compound (5) (20 ppm)	80	64	76	13	
Compound (6) (0.2 ppm)	92	67	93	7	
Compound (7) (0.2 ppm)	91	79	90	22	

#### Example 7

Isolation of Magnaporthe grisea HIK1 gene

[0107] Total RNA was prepared from Magnaporthe grisea. 100 mg of a hypha of Magnaporthe grisea P-37 strain which had been grown on a potato dextrose agar medium (PDA medium manufactured by NISSUI Pharmaceutical Co., Ltd.) was scratched off, and this was ground using a mortar and a pestle in liquid nitrogen. A RNA was prepared from frozen ground powder using RNeasy Plant Mini Kit (QIAGEN). A frozen ground powder together with liquid nitrogen was transformed to a 50 ml sample tube and, after liquid nitrogen was all volatilized off, a solution obtained by adding 10  $\mu$ L of mercaptoethanol was added per 1 ml of a buf fer RLC attached to the kit was added, followed by stirring. Further, after ground powder was well dispersed by a few pipettings, a temperature was maintained at 56°C for 3

minutes. Thereafter, a solution containing ground powder was supplied to QIAshredder spin column attached to the kit, and centrifuged at 8,  $000\times g$  for 2 minutes. The filtration supernatant was transferred to a fresh sample tube, a 0.5-fold volume of 99.5% ethanol was added, and the material was well mixed by pipetting. This mixtured solution was supplied to RNeasy mini spin column attached to the kit, and centrifuged at  $8,000\times g$  for 1 minute. The filtrate was discarded, 700  $\mu$ L of Buffer RW1 attached to the kit was added, centrifuged at  $8,000\times g$  for 1 minute, and the filtrate was discarded. Further, the residue was added 500  $\mu$ L of Buffer RPE attached to the kit, and centrifuged at  $8,000\times g$  for 1 minute, and the filtrate was discarded. This procedure was repeated twice. Finally, an upper filter part was transferred to a fresh sample tube, supplied 30  $\mu$ L of RNase-free sterilized water, and centrifuged at  $8,000\times g$  for 1 minute, and total RNA was dissolved into the filtrate. This dissolution procedure was repeated twice.

[0108] Then, a cDNA was synthesized using ThermoScript RT-PCR System (Invitrogen) while using total RNA as a template. A solution in which 9.0  $\mu$ L of total RNA was mixed into 1.0  $\mu$ L of 50 mM Oligo(dt)<sub>20</sub> attached to the kit and 2.0  $\mu$ L of 10 mM dNTP Mix was treated at 65°C for 5 minutes, and rapidly cooled on ice. To this solution were added 4  $\mu$ L of 5×cDNA Synthesis Buffer attached to the kit, 1  $\mu$ L of 0.1M DTT, 1  $\mu$ L of RNase OUT, 1  $\mu$ L of ThermoScript RT and 1  $\mu$ L of sterilized distilled water, to react them at 50°C for 60 minutes and, thereafter, the reaction was stopped by heating treatment at 85°C for 5 minutes. Further, 1  $\mu$ L of RNaseH attached to the kit was added to this reaction solution, the materials were reacted at 37°C for 20 minutes, and a RNA as a template was degraded to obtain a cDNA.

[0109] A DNA having a nucleotide sequence encoding an amino acid sequence of Magnaporthe grisea HIK1 (hereinafter, referred to as HIK1 DNA in some cases) was amplified by PCR using this cDNA as a template. A PCR was performed using an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 18 and an oligonucleotide consisting of the nucleotide sequence represented by SEQ IDNO: 19, to amplify a DNA having the nucleotide sequence represented by SEQ IDNO: 17. The PCR was performed using KOD-Plus-(TOYOBO) under the amplifying conditions that a temperature was maintained at 94°C for 2 minutes and, thereafter, 35 cycles of incubation were repeated, each cycle comprising maintaining a temperature at 94°C for 15 seconds, then, at 55°C for 30 seconds, then, at 68°C for 6 minutes. The FCR reaction solution (50  $\mu$ L) was prepared by adding 2  $\mu$ L of the aforementioned cDNA, 5  $\mu$ L of 10×Buffer, 5  $\mu$ L of 2 mM dDNPs, 2  $\mu$ L of 25 mM MgSO<sub>4</sub>, each 1  $\mu$ L of 10  $\mu$ M oligonucleotide primers, 33  $\mu$ L of sterilized distilled water and 1  $\mu$ L of KOD-Plus-. After the reaction, a part of the reaction solution was separated with 1.0% agarose gel electrophoresis, and stained with ethidium bromide. It was confirmed that about 4 kb of the DNA (HIK1 DNA) was amplified.

## 30 Example 8

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Construction of an expression plasmid of Magnaporthe grisea HIK1 gene and preparation of transformed budding yeast.

[0110] The HIK1 DNA was cloned into a cloning vector pBluesripit SK II (+). About 4 kb of the aforementioned DNA (HIK1 DNA) was purified from the PCR reaction solution prepared in Example 7 using QIAquick PCR Purification Kit -(QIAGEN) according to the attached manual. About 4 kb of the purified DNA (HIK1 DNA) was digested with restriction enzymes of Spel and HindIII and, on the other hand, after the cloning vector pBluescript SK II (t) (manufactured by Stratagene) was also digested with restriction enzymes Spel and HindIII, each of which was separated with 1.0% agarose gel electrophoresis, and a part of the gel containing a desired DNA was excised. The NIK1 DNA digested with Spel and HindIII and the cloning vector digested with Spel and HindIII were recovered form the gel using QIAquick Gel Extraction Kit (QIAGEN) according to the attached manual. The HIK1 DNA was inserted between Spel site and HindIII site in the multicloning site of the cloning vector using Ligation Kit Ver.2 (TaKaRa) according to the attached manual, to construct a plasmid pBlueHIK1. A nucleotide sequence of the resulting plasmid was analyzed with a DNA sequencer (Model 3100, Applied Biosystems) after a sequencing reaction using BigDye terminator v3.0 Cycle Sequence FS Ready Reaction Kit (Applied Biosystems) according to the attached manual. The sequencing reaction was performed using an oligonucleotide consisting of the nucleotide sequence represented by any of SEQ ID NOs.20 to 29 as a primer under the amplifying conditions that 35 cycles of incubation were repeated, each cycle comprising maintaining a temperature at 96°C for 10 seconds, then, at 50°C for 5 seconds, then, at 60°C for 2 minutes. As a result, the nucleotide sequence represented by SEQ ID NO: 17 was obtained, and it was confirmed that the plasmid pBlueHIK1 harbored a DNA having a nucleotide sequence encoding an amino acid sequence of HIK1.

[0111] Then, the HIK1 DNA was inserted into a shuttle vector p415ADH (ATCC87312) replicable in yeast and Escherichia coli. The plasmid pBlueHIK1 prepared as described above was digested with restriction enzymes Spel and HindIII and, on the other hand, after the shuttle vector p415ADH (ATCC87312) was also digested with restriction enzymes Spel and HindIII, each of which was separated with 1.0% agarose gel electrophoresis, and a part of the gel containing a desired DNA was excised. The HIK1 DNA digested with Spel and HindIII and the shuttle vector digested with Spel and HindIII were recovered from the gel using QIAquick Gel Extraction Kit (QIAGEN) according to the attached manual. The HIK1 DNA was inserted between Spel site and HindIII site in the multicloning site of the shuttle vector using Ligation Kit Ver.2 (TaKaRa) according to the attachedmanual, to construct an expression plasmid pADHHIK1.

[0112] The prepared expressed plasmid pADHHIK1 was introduced into budding yeast (Saccharomyces cerevisiae) AH22 strain (IFO10144 andTM182 strain (MaedaT. et al. (1994) Nature vol. 369, pp242-245) according to the method described in Geitz RD & Woods RA (1994) Molecular Genetics of Yeast: Practical Approaches ed. Johnson JA, Oxford University Press pp124-134. By utilizing disappearance of leucine auxotrophy in the resulting transformed budding yeast, the transformed budding yeast AH22 strain (AH21-HIK1) was selected on a Glu-Leu agar medium, and the transformed budding yeast TM182 strain (TM182-HIK1) was selected on a Glu-Ura-Leu agar medium. It was confirmed that the resulting TM182-HIK1 grows even transferred to a Glu-Ura-Leu medium.

#### Example 9

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Antifungal compound sensitivity test of transformed budding yeast TM182-HIK1.

[0113] The transformed budding yeast AH22-HIK1 prepared in Example 8 was cultured while shaking at 30°C for 24 hours in a Glu-Leu medium. As a control, the AH22 strain was similarly cultured while shaking at 30°C for 24 hours in a Glu medium. The absorbance at 600 nm of a cell suspension of each of the grown transformed budding yeasts was measured, and a cell suspension diluted with each medium to the absorbance of 0.1 was prepared. Further, a cell suspension in which the aforementioned cell suspension of the transformed budding yeast AH22-HIK1 was diluted 50-fold with a Glu-Leu medium, and a cell suspension in which the aforementioned cell suspension of AH22 strain was diluted 50-fold with a Glu medium were prepared. A suspension in which each of compounds (1) to (3) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 200 ppm, a solution in which each of Compounds (4) to (5) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 600 ppm, and a solution in which each of Compounds (6) and (7) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 20 ppm were prepared, and two microplates were prepared in which each 1.0 μL per well of the Compound solution and DMSO as a control were dispensed into two wells. In one microplate among them, each 100 µL of a cell suspension of the transformed budding yeast AH22-hiki which had been prepared by dilution as described above was dispensed, and cultured by allowing to stand at 30°C for 23 hours. In another microplate, each 100 µL of the cell suspensions of control yeast AH22 strain which had been prepared by dilution as described above was dispensed, and cultured by allowing to stand at 30°C for 27 hours. After culturing, the absorbance at 600 nm of each well was measured with a microplate reader.

[0114] Similarly, the transformed budding yeast TM182-HIK1 prepared in Example 8 was cultured at 30°C for 24 hours in a Glu-Ura-Leumedium. The absorbance at 600 nmof a cell suspension of the grown transformed budding yeast was measured, and a cell suspension diluted with each medium to the absorbance of 0.1 was prepared. Further, a cell suspension in which the aforementioned cell suspension of the transformed budding yeast TM182-HIK1 was diluted 50-fold with a Glu-Ura-Leu medium and, as a control, a cell suspension in which the suspension was diluted 50-fold with a Glu-Ura-Leu medium were prepared. A suspension in which each of Compounds (1) to (3) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 200 ppm, a solution in which each of Compounds (4) and (5) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 600 ppm, and a solution in which each of Compounds (6) and (7) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 20 ppm were prepared, and two microplates were prepared in which each 1.0 µL per well of the Compound DMSO solution and DMSO as a control were dispensed into two wells. In one microplate among them, each 100 µL of cell suspensions of the transformed budding yeast TM182-HIK1 which had been prepared by dilution with a Glu-Ura-Leu medium as described above was dispensed, and cultured by allowing to stand at 30°C for 27 hours. In another microplate, as described above, as a control, each 100 μL of cell suspensions of the transformed budding yeast TM182-HIK1 which had been prepared by dilution with a Gal-Ura-Leu medium was dispensed, and cultured by allowing to stand at 30°C for 27 hours. After culturing, the absorbance at 600 nmof each well was measured with a microplate reader.

[0115] Degree of growths of both of the transformed budding yeasts cultured in the presence of Compounds (1) to (7) and budding yeast as a control therefor are shown in Table 3. Degrees of growths of both of the transformed budding yeasts and budding yeast as a control thereofor are shown by a relative value in percentage, letting the absorbance of 600 nm at the concentration of the Compound of 0 ppm to be 100. It was confirmed that an inhibiting degree of growth of TM182-HIK1 by each test substance was greater than an inhibiting degree of growth of AH22-HIK1 by each test substance, and the TM182-HIK1 was a transformed cell with the enhanced sensitivity to an antifungal compound as compared with AH22-HIK1.

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Table 3

Test substance (final concentration)	Degree of growth of budding yeast (%)			
	AH22	AH22-HI-K1 TM182-HIK1 Glu-Leu medium Gal-Ura-Leu medium		
	Glu medium		Gal-Ura-Leu medium	Glu-Ura-Leu medium
Compound (1) (2.0 ppm)	85	89	100	62
Compound (2) (2.0 ppm)	96	84	94	79
Compound (3) (2.0 ppm)	99	104	100	30
Compound (4) (6.0 ppm)	97	92	97	63
Compound (5) (6.0ppm)	93	99	106	22
Compound (6) (0.2 ppm)	101	98	104	11
Compound (7) (0.2 ppm)	89	102	87	9

Example 10

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Amplification of osmosensitivie histidine kinase gene fragment from other filamentas fungus

- (1) Preparation of Total RNA of Fusarium oxysporum
- [0116] Total RNA was prepared from Fusarium oxysporum. 100 mg of a hypha of Fusarium oxyporum RJN1 strain grown on a potato dextrose agarose medium (PDA medium, manufactured by NISSUI Pharmaceutical Co., Ltd.) was collected, and this was ground using a mortar and a pestle in liquid nitrogen. Total RNA was prepared from frozen ground powder using RNeasy Plant Mini Kit (QIAGEN) according to the method described in Example 1.
- (2) Preparation of Total RNA of Mycospharella tritici
- [0117] Total RNA was prepared from Mycospharella tritici. Spore of Mycospharella tritici St-8 strain grown on a potato dextrose agarose medium (PDA medium, manufactured by NISSUI Pharmaceutical Co., Ltd.) was added to 100 ml of PD broth (DIFCO), and this was cultured at 20°C and 150rpm for 4 days using a 500 ml volume Erlenmeyer flask. 8 ml of the culture solution was centrifuged to remove the supernatant, and 300 mg of a wet weight of cells were transferred to a mortar and ground in liquid nitrogen using a pestle. Total RNA was prepared from frozen ground powder according to the method described in Example 1.
- (3) Preparation of total RNA of Thanatephorus cucumeris
- [0118] Total RNA was prepared from Thanatephorus cucumeris. Hypha of Thanatephorus cucumeris Rs-18 strain grown on a potato dextrose agar medium (PDA medium, manufactured by NISSUI Pharmaceutical Co., Ltd.) was added to 100 ml of PD broth (DIFCO), and cultured by allowing to stand at 25°C for 4 days using a 500 ml volume Erlenmeyer flask. 8 ml of the culture solution was centrifuged to remove the supernatant, 300 mg of a wet weight of hypha were transferred to a mortar, and ground in liquid nitrogen using a pestle. Total RNA was prepared from frozen ground powder using Rneasy Plant Mini Kit (QIAGEN) according to the method described in Example 1.
- (4) Preparation of total RNA of Phytophthora infestans
- [0119] Total RNA was prepared from Phytophthora infestans. Hypha of Phytophthora infestans Pi-5 strain grown on a rye agar medium (rye 60g, sucrose 15g, agar 20g/1L) was added to 20 ml of a rye medium (rye 60g, sucrose 15g/1L), and cultured at 20°C and 150rpm for 5 days using a 300 ml of volume Erlenmeyer flask. 20 ml of the culture solution was centrifuged to remove the supernatant, a wet weight of 200 mg of cells were transferred to a mortar, and ground using a pestle in liquid nitrogen. Total RNA was prepared from frozen ground powder using RNeasy Plant Mini Kit (QIAGEN) according to the method described in Example 1.

(5) Amplification of osmosensing histidine kinase gene fragment by PCR

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[0120] Using the total RNA of Magnaporthe grisea prepared in Example 7, the total RNA of Fusariumoxysporumprepared in Example 10 (1), the total RNA of Mycospharella tritici prepared in Example 10 (2), the total RNA of Thanatephorus cucumeris prepared in Example 10 (3), or the total RNA of Phytophthora infestans prepared in Example 10 (4), amplification of a DNA having a nucleotide sequence encoding a part of an amino acid sequence of osmosensing histidine kinase was performed.

[0121] First, a cDNA was synthesized using ThermoScript RT-PCR System (Invitrogen) and using each of total RNAs as a template. A solution in which 4.0  $\mu$ L of each of total RNAs and 5.0  $\mu$ L of sterilized distilled water were mixed into 1.0  $\mu$ L of 50 mM Oligo (dT)<sub>20</sub> attached to the kit and 2.0  $\mu$ L of 10 mM dNTP Mix was prepared, and a cDNA was synthesized according to the method described in Example 1.

[0122] A PCR was performed using each cDNA as a template. As primers, a primer pair shown in Table 4 was used. A size of a DNA which is predicted to be amplified by PCR using each primer pair based on the nucleotide sequence represented by SEQ ID NO: 2 is shown in Table 4.

Table 4

	Table 4	
Primer	Primer	DNA to be amplified
SEQ ID NO: 30	SEQ ID NO: 35	368bp
SEQ ID NO: 30	SEQ ID NO: 36	374bp
SEQ ID NO 30	SEQ ID NO: 37	383bp
SEQ ID NO: 31	SEQ ID NO: 35	359bp
SEQ ID NO: 31	SEQ ID NO: 36	365bp
SEQ ID NO: 31	SEQ ID NO: 37	374bp
SEQ ID NO: 32	SEQ ID NO: 38	3019bp
SEQ ID NO: 32	SEQ ID NO: 40	3052bp
SEQ ID NO: 33	SEQ ID NO: 38	2927bp
SEQ ID NO: 33	SEQ ID NO: 40	2960bp
SEQ ID NO: 34	SEQ ID NO: 38	2867bp
SEQ ID NO: 34	SEQ ID NO: 40	2900bp
SEQ ID NO: 30	SEQ ID NO: 39	1424bp
SEQ ID NO: 30	SEQ ID NO: 40	1442bp
SEQ ID NO: 31	SEQ ID NO: 39	1415bp
SEQ ID NO: 31	SEQ ID NO: 40	1433bp
	SEQ ID NO: 30 SEQ ID NO: 30 SEQ ID NO: 30 SEQ ID NO: 31 SEQ ID NO: 31 SEQ ID NO: 31 SEQ ID NO: 32 SEQ ID NO: 32 SEQ ID NO: 33 SEQ ID NO: 33 SEQ ID NO: 34 SEQ ID NO: 30 SEQ ID NO: 30 SEQ ID NO: 30	Primer         Primer           SEQ ID NO: 30         SEQ ID NO: 35           SEQ ID NO: 30         SEQ ID NO: 36           SEQ ID NO: 30         SEQ ID NO: 37           SEQ ID NO: 31         SEQ ID NO: 35           SEQ ID NO: 31         SEQ ID NO: 36           SEQ ID NO: 31         SEQ ID NO: 37           SEQ ID NO: 32         SEQ ID NO: 38           SEQ ID NO: 32         SEQ ID NO: 40           SEQ ID NO: 33         SEQ ID NO: 38           SEQ ID NO: 34         SEQ ID NO: 38           SEQ ID NO: 34         SEQ ID NO: 38           SEQ ID NO: 34         SEQ ID NO: 40           SEQ ID NO: 30         SEQ ID NO: 40           SEQ ID NO: 30         SEQ ID NO: 40           SEQ ID NO: 31         SEQ ID NO: 40           SEQ ID NO: 31         SEQ ID NO: 40

[0123] A PCR was performed using KOD-Plus-(TOYOBO) under the amplifying conditions that a temperature was maintained at 94°C for 2 minutes and, thereafter, 35 cycles of incubation were repeated, each cycle comprising maintaining a temperature at 94°C for 15 seconds, then, at 55°C for 30 seconds further, at 68°C for 1 minutes. When primer pairs 1 to 6 were used, the incubation at 68°C in the cycle was for 1 minutes. When the primer pairs 7 to 12 were used, the incubation at 68°C in the cycle was for 5 minutes. When the primer pairs 13 to 16 were used, the incubation at 68°C in the cycle was for 3 minutes. The PCR reaction solution (25  $\mu$ L) was prepared by adding 0.5  $\mu$ L of the cDNA, 2.5  $\mu$ L of 10xbuffer, 2.5  $\mu$ L of 8 mM dNTPs, 1.0  $\mu$ L of 25 mM MgSO<sub>4</sub>, each 0.5  $\mu$ L of 10  $\mu$ M oligonucleotide primers, 17  $\mu$ L of sterilized distilled water and 0.5  $\mu$ L of KOD-Plus-. The PCR reaction solution after the reaction was analyzed with 1% or 4% agarose gel electrophoresis.

[0124] When primer pairs 1, 2, 3, 4, 5 or 6 were used and a cDNA of Magnaporthe grisea was used as a template, amplification of predicted size of DNA was observed. When primer pairs 2, 3,, 7, 8, 9, 10, 11 or 12 were used, and a cDNA of Fusarium oxysporum was used as a template, amplification of a predicted size of DNA was observed. When the primer pairs 3, 5, 6, 13, 14, 15 or 16 were used, and cDNA of Mycospharella Tritici was used as a template, amplification of predicted size of DNA was observed. When primer pairs 2, 3, 5 or 6 were used, and cDNA of Thanate-phorus cucumeris was used as a template, amplification of a predicted size of a DNA was observed. When the primer pairs 5 or 6 were used, and cDNA of Phytophthora infestans was used as a template, amplification of predicted size

of DNA was observed.

Example 11

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- 5 Isolation of Fusarium oxysporum FoOS-1 gene
  - (1) Analysis of Fusarium oxysporum FoOS-1 gene fragment
  - [0125] The amplified DNA was purified from the reaction solution of PCR which had been performed by using a cDNA of Fusarium oxysporum as a template and using a primer pair 9 in Example 10 (5), using QIAquick PCR Purification Kit (QIAGEN) according to the attached instruction.
  - [0126] Adenine was added to the 3'-terminal of the purified DNA using Ex-Taq (TaKaRa) (hereinafter, referred to as 3' A addition). The reaction solution (20  $\mu$ L) for 3' A addition was prepared by adding 15.3  $\mu$ L of a solution of the aforementioned purified DNA, 2.0  $\mu$ L of 10 × buffer, 2.5  $\mu$ L of 10 mM dNPTs and 0.2  $\mu$ L of Ex Taq, and this was maintained at 72°C for 30 minutes.
  - [0127] Thus the 3' A-added DNA and the pCR2. 1-TOPO cloning vector (Invitrogen) were ligated according to the instruction attached to the cloning vector, after that, which was introduced into Escherichia coli JM109 (TaKaRa). A plasmid DNA was purified from the resulted Escherichia coli transformant using QIAprep Spin Miniprep Kit (QIAGEN). A nucleotide sequence of the plasmid DNA was analyzed with a DNA sequencer (Model 3100, Applied Biosystems) after a sequencing reaction employing the resulting plasmid DNA as a template, and using an oligonucleotide consisting of the nucleotide sequence represented by any of SEQ IDNOs: 28, 29, and 45 to 48 as a primer, and using BigDye Terminator v3.0 Cycle Sequencing Ready Reaction Kit (Applied Biosystems Japan) according to the instruction attached to the kit. The sequencing reaction was performed under the amplifying conditions that 35 cycles of incubation were repeated., each cycle comprising maintaining a temperature at 96°C for 10 seconds, then, at 50 °C for 5 seconds, further, at 60°C for 2 minutes. As a result, a nucleotide sequence represented by base numbers 663 to 3534 of the nucleotide sequence represented by SEQ ID NO: 42 was read.
  - (2) Analysis of full length FoOS-1 gene of Fusarium oxysporum
- [0128] A DNA having a nucleotide sequence extending toward to the 5' upstream region from a nucleotide number 663 of the nucleotide sequence represented by SEQ ID Mo. 42 was cloned using SMART RACE cDNA Amplification Kit (CLONTECH) according to the instruction attached to the kit. 1.0 μL of CDS-primer attached to the kit, and 1.0 μL of SMART IIA Oligo were mixed into 3 μL (230ng) of the total RNA prepared in Example 10 (1) to prepare a reaction solution. The reation solution was maintained at 70°C for 2 minutes and maintained on ice for 2 minutes. To the reaction 35 solution were added 2  $\mu$ L of 5 imes First-Strand buffer attached to the kit, 1  $\mu$ L of 20 mM DTT, 1  $\mu$ L of 10 mM dNPT Mix -and 1 μL of PowerScript Reverse Transcriptase and mixed, and the mixture was maintained at 42°C for 1. 5 hours. To the reaction solution after temperature maintenance was added 100 µL of Tricine-EDTA buffer attached to the kit, and a temperature was maintained at 72°C for 7 minutes to prepare 5' RACE ready cDNA. PCR amplifying 5' upstream region was performed by using this 5' RACE ready cDNA as a template. A PCR reaction solution was obtained by adding  $5.0 \,\mu\text{L}$  of  $10 \times$  Advantage 2 buffer,  $1.0 \,\mu\text{L}$  of  $10 \,\mu\text{M}$  dNTP Mix and  $1.0 \,\mu\text{L}$  of  $50 \times$  Advantage 2 Polymerase Mix attached to the kit to 2.5  $\mu$ L of 5' RACE ready cDNA and mixing them, and adding 5. 0  $\mu$ L of 10 $\times$ Universal Primer A Mix attached to the kit as a primer, and 1.0 μL of a 10 μM solution of an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 43, and adding sterilized distilled water to a total amount of 50  $\mu$ L. This reaction solution was subjected to repetition of 5 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 5 seconds, then, at 72°C for 2 minutes, further repetition of 5 cycles of incubation, each cycle comprising a maintaining a temperature at 94°C for 5 seconds, then, at 70°C for 10 seconds, then, at 72°C for 2 minutes, further repetition of 25 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 5 seconds, then, at 68°C for 10 seconds, then, at 72°C for 2 minutes, followed by maintaining a temperature at 72°C for 7 minutes. The PCR reaction solution and the pCR2.1-TOPO cloning vector (Invitrogen) were ligated according to the instruction attached to the 50 cloning vector, after that, which was introduced into Escherichia coli JM109 (TaKaRa). A plasmid DNA was purified from the resulting Escherichia coli transformant using QIAprep Spin Miniprep Kit (QIAGEN). A nucleotide sequence was analyzed using the resulting plasmid DNA as a template, and using aprimer consisting of the nucleotide sequence represented by any of SEQ ID NOs: 29, 49 and 54 according to the method described in Example 11 (1). As a result, a nucleotide sequence represented by nucleotide numbers 1 to 662 of the nucleotide sequence represented by SEQ
  - [0129] Further, a DNA having a nucleotide sequence extending toward to the 3' downstream region from nucleotide number 3534 of the nucleotide sequence represented by SEQ ID NO: 42 was cloned. 1. 0  $\mu$ L of CDS-primer attached to the kit and 1.0  $\mu$ L of sterilized distilled water were mixed into 3  $\mu$ L (230ng) of the total RNA prepared in Example 10

- (1), the mixture was maintained at 70°C for 2 minutes, and maintained on ice for 2 minutes. 3' RACE ready cDNA was prepared using the reaction solution as in preparation of 5' RACE ready cDNA. PCR amplifying 3' downstream region was performed using this 3' RACE ready cDNA as a template. A PCR reaction solution was prepared by mixing 5.0  $\mu L$  of 10 $\times$  Advantage 2 buffer attached to the kit, 1.0  $\mu L$  of 10 mM dNTP Mix and 1.0  $\mu L$  of 50 $\times$  Advantage 2 Polymerase Mix into 2.5 μL of 3' RACE ready cDNA, adding 5.0 μL of 10×Universal Primer A Mix attached to the kit as a primer, and 1. 0 µL of a 10 µM solution of an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 42, and adding sterilized distilled water to a total amount of 50 µL. This reaction solution was subjected to repetition of 5 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 5 seconds, then, at 72°C for 2 minutes, further repetition of 5 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 5 seconds, then, at 70°C for 10 seconds, then, at 72°C for 2 seconds, further repetition of 25 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 5 seconds, then, at 68°C for 10 seconds, then, at 72°C for 2 minutes, followed by maintaining a temperature at 72°C for 7 minutes. The PCR reaction solution and the pCR2.1-TOPO cloning vector (Invitrogen) were ligated to the vector according to the instruction attached to the kit, after that, which was introduced into Escherichia coli JM109 (TaKaRa). Aplasmid DNAwas purified from the resulting Escherichia coli transformant using QIAprep Spin Miniprep Kit (QIAGEN). A nucleotide sequence was analyzed, using the resulting plasmid DNA as a template, and using a primer consisting of the nucleotide sequence represented by any of SEQ ID NOs: 29, 50 and 54, according to the method described in Example 11 (1). As a result, a nucleotide sequence represented by nucleotide numbers 3535 to 3882 of the nucleotide sequence represented by SEQ ID NO: 42 was read.
- [0130] All analyzed nucleotide sequences were joined and, as a result, the nucleotide sequence represented by SEQ ID NO: 42 was obtained. The nucleotide sequence represented by SEQ ID NO: 42 consists of 3882 bases (including termination codon), and was a nucleotide sequence encoding 1293 amino acid residues (SEQ ID NO: 41). A molecular weight of a protein having the amino acid sequence represented by SEQ ID NO: 41 was calculated to be 141818 Da.
- 25 (3) Isolation of full length Fusarium oxysporum FoOS1 gene
  - [0131] A DNA having a nucleotide sequence encoding an amino acid sequence of Fusarium oxysporum FoOS1 (hereinafter, referred to as FoOS-1 DNA in some cases) was amplified by PCR using the 5' RACE ready cDNA prepared in Example 11 (2) as a template. By performing a PCR using, as a primer, an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 52 and an oligonucleotide consisting of the nucleotide sequence represented by SEQ IDNO: 42 was amplified. The PCR was performed using KOD-Plus- (TOYOBO) under the amplifying conditions that a temperature was maintained at 94°C for 2 minutes and, thereafter, 35 cycles of incubation were repeated, each cycle comprising maintaining a temperature at 94°C for 15 seconds, then, at 55°C for 30 seconds, further, at 68°C for 6 minutes. The PCR reaction solution (50 μL) was prepared by adding 2.5 μL of 5' a RACE ready cDNA, 5.0 μL of 10×buffer, 5.0 μL of 2 mM dNTPs, 2.0 μL-of 25 mM MgSO<sub>4</sub>, each 1.0 μL of 10 μM oligonucleotide primers, 32.5 μL of sterilized distilled water and 1.0 μL of KOD Plus-. After the reaction, a part of the PCR reaction solution was separated by 1% agarose gel electrophoresis, and stained with ethidium bromide. It was confirmed that about 4 kb of the DNA (FoOS1 DNA) was amplified.
- 40 Example 12

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 $Construction\ of\ expression\ plasmid\ of\ Fusarium\ oxysporum\ FoOS1\ gene\ and\ preparation\ of\ transformed\ budding\ yeast$ 

- [0132] The FoOS1 DNA was cloned into a pCR2 .1-TOPO cloning vector (Invitrogen). About 4 kb of the DNA (FoOS-1 DNA) was purified from the PCR reaction solution prepared in Example 11 (3) using QIAquick PCR Purification Kit (QIAGEN) according to the instruction attached to the kit. 3'A addition was performed on about 4 kb of the purified DNA (FoOS-1 DNA) according to the method described in Example 11 (1). The 3'A-added about 4 kb DNA (FoOS-1 DNA) and the pCR2.1-TOPO cloning vector (Invitrogen) were ligated according to the manual attached to the cloning vector to construct a plasmid pCRFoOSI. A nucleotide sequence of the resulting plasmid was analyzed according to the method described in Example 11(1). As a primer, an oligonucleotide consisting of the nucleotide sequence represented by any of SEQ ID NOs: 29, 43 to 51, and 54 was used. As a result, the nucleotide sequence represented by SEQ ID NO: 42 was obtained, and it was confirmed that the plasmid pCRFoOS1 was a plasmid containing the FoOS-1 DNA.
- [0133] The FoOS-1 DNA contained in the thus prepared plasmid pCRFoOS1 was cloned into a shuttle vector p415ADH replicable in yeastand Escherichia coli to construct an expression plasmid. The plasmid pCRFoOS1 was digested with restriction enzymes Spel and Pstl and, on the other hand, the shuttle vector p415ADH was also digested with restriction enzymes Spel and Pstl. Each of them was separated by 0.8% agarose gel electrophoresis, a part of the gel containing the FoOS-1 DNA digested with Spel and Pstl and the shuttle vector p415ADH digested with Spel

and PstI was excised, and the FoOS-1 DNA and the shuttle vector were recovered from the gel using QIAquick Gel Extraction Kit (QIAGEN) according to the attached manual. The FoOS-1 DNA was inserted between SpeI site and PstI site in the multicloning site of the shuttle vector using Ligation Kit Ver.2 (TaKaRa) according to the manual attached to the kit, whereby, an expression plasmid pADHFoOS1 was constructed. A nucleotide sequence of the resulting expression plasmid was analyzed according to the method described in Example 11 (1). As a primer, an oligonucleotide consisting of the nucleotide sequence represented by any of SEQ ID NO: 43 to 53 was used. As a result, the nucleotide sequence represented by SEQ ID NO: 42 was obtained, and it was confirmed that the expression plasmid pADHFoOS1 harbored a DNA having a nucleotide sequence encoding an amino acid sequence of FoOS-1.

[0134] The prepared expression plasmid pADHFoOS1 was introduced into budding yeast AH22 strain and TM182 strain according to the method described in Example 2. By utilizing the disappearance of leucine auxotrophy in the resulting transformed budding yeast, the transformed budding yeast AH22 strain (AH22-FoOS1) was selected on a Glu-Leu agar medium, and the transformed budding yeast TM182 strain (TM182-FoOS1) was selected on a Gal-Ura-Leu agar medium. It was confirmed that the resulting TM182-FoOS1 grows even when transplanted to a Glu-Ura-Leu medium.

#### Example 13

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Antifungal compound sensitivity test of transformed budding yeast TM182-FoOS1

[0135] The transformed budding yeast AH22-FoOS1 prepared in Example 12 was cultured while shaking at 30°C for 18 hours in a Glu-Leu medium. As a control, the AH22 strain was similarly cultured while shaking at 30°C for 18 hours in a Glu medium. The absorbance at 600 nm of each grown transformed budding yeast in a cell suspension was measured, and cell suspension diluted with sterilized distilled water to the absorbance of 0.1 was prepared. Further, a cell suspension in which the aforementioned cell suspension of the transformed budding yeast AH22-FoOS1 was diluted 50-fold with a Glu-Leu medium, and a cell suspension in which the aforementioned cell suspension of the AH22 strain was diluted 50-fold with a Glu medium were prepared.

[0136] A solution in which each of Compounds (1) to (3) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 600 ppm, a solution in which each of Compounds (4) and (5) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 2000 ppm, and a solution in which Compounds (6) and (7) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 20 ppm were prepared, and two microplates were prepared in which each 1.0  $\mu$ L per well of the Compound solution and DMSO as a control were dispensed. In one microplate among them, each 100  $\mu$ L of cell suspensions of the transformed budding yeast AH22-FoOS1 which had been prepared by dilution as described above was dispensed, and cultured by allowing to stand at 30°C for 26.5 hours. In another microplate, each 100  $\mu$ L of cell suspensions of the control yeast AH22 strain which had been prepared by dilution as described above was dispensed, and cultured by allowing to stand at 30°C for 24.5 hours. After culturing, the absorbance at 600 nm of each well was measured with a microplate reader.

[0137] Similarly, the transformed budding yeast TM182-FoOS1 prepared in Example 12 was cultured at 30°C for 18 hours in a Glu-Ura-Leu medium. The absorbance at 600 nm of the grown transformed budding yeast in a cell suspension was measured, and a cell suspension diluted with sterilized distilled water to the absorbance of 0.1 was prepared. Further, a cell suspension in which the transformed budding yeast TM182-FoOS1 was diluted 50-fold with a Glu-Ura-Leu medium and, as a control, a cell suspension in which the yeast was diluted 50-fold with a Gal-Ura-Leu medium were prepared.

[0138] A solution in which each of Compounds (1) to (3) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 600 ppm, a solution in which each of Compounds (4) and (5) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 2000 ppm, and a solution in which each of Compounds (6) and (7) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 20 ppm were prepared, and two microplates were prepared in which each 2.0  $\mu$ L per well of the Compound-DMSO solution and DMSO as a control were dispensed into 2 wells. In one microplate among them, each 100  $\mu$ L of cell suspensions of the transformed budding yeast TM182-FoOS1 which had been prepared by dilution with a Glu-Ura-Leu medium as described above was dispensed, and cultured by allowing to stand at 30°C for 25 hours. In another microplate, as described above, as a control, each 100  $\mu$ L of cell suspensions of the transformed budding yeast TM182-FoOS1 which had been prepared by dilution with a Gal-Ura-Leu medium was dispensed, and cultured at 30°C for 51 hours. After culturing, the absorbance at 600 nm of each well was measured with a microplate reader.

[0139] A degree of growth of each transformed budding yeast cultured in the presence of Compounds (1) to (7) is shown in Table 5. A degree of growth of the transformed budding yeast is expressed as a relative value in percentage, letting the absorbance at 600 nm at the concentration of the Compound of 0 ppm to be 100. It was confirmed that an inhibiting degree of growth of the transformed budding yeast TM182-FoOS1 by each test substance was grater than an inhibiting degree of growth of the transformed budding yeast AH22-FoOS1 by each test substance, and the transformed budding yeast AH22-FoOS1 by each test substance, and the transformed budding yeast AH22-FoOS1 by each test substance.

formed budding yeast TM192-FoOS1 was a transformed cell with the enhanced sensitivity to an antifungal compound as compared with the transformed budding yeast AH22-FoOS1.

Table 5

Test substance (final concentration)	Degree of growth of budding yeast			
	AH22	AH22-Fo OS1	TM182-FoOS1	
	Glu medium	Glu-Leu medium	Gal-Ura-Leu medium	Gul-Ura-Leu medium
Compoud (1) (6 ppm)	88	81	116	26
Compoud (2) (6 ppm)	91	91	87	55
Compoud (3) (6 ppm)	87	86	99	22
Compoud (4) (20 ppm)	96	90	104	20
Compoud (5) (20 ppm)	80	71	80	57
Compoud (6) (0.2 ppm)	92	69	99	7
Compoud (7) (0.2 ppm)	91	88	89	21

Example 14

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Isolation of Mycospharella tritici StOS-1 gene

(1) Analysis of Mycospharella tritici StOS-1 gene fragment

[0140] The amplified DNA was purified from the reaction solution of PCR which had been performed using a primer pair 16 and using a cDNA of Mycospharella tritici as a template in Example 10 (4), using QIAquick PCR Purification Kit (QIAGEN) according to the instruction attached to the kit. 3'A addition was performed on the purified DNA according to the method described in Example 11 (1). The 3'A-added DNA and the pCR2.1-TOPO cloning vector (Invitrogen) were ligated according to the instruction attached to the cloning vector, and this was introduced into Escherichia coli JM109 (TaKaRa).

[0141] DNA was purified from the resulting Escherichia coli transformant by colony PCR using Ex Taq HS (TaKaRa). The PCR reaction solution (15  $\mu$ L) was prepared by mixing 1.5  $\mu$ L of 10×buffer, 2.25  $\mu$ L of 10 mM dNTPs, 0.15  $\mu$ L of Ex Taq HS, each 0.4  $\mu$ L of a 10  $\mu$ M solution of an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 66 and a 10  $\mu$ M solution of an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 67, and 10. 3  $\mu$ L of sterilized distilled water, and adding a part of the Escherichia coli transformant colony thereto. PCR was performed under the amplifying conditions that this reaction solution was maintained at 97°C for 2 minutes and, thereafter, 35 cycles of incubation were repeated, each cycle comprising maintaining a temperature at 97°C for 15 seconds, then, at 55°C for 15 seconds, then, at 72°C for 3 minutes. The amplified DNA was purified from the PCR reaction solution after temperature maintenance using QIAquick PCR Purification Kit (QIAGEN) according to the instruction attached to the kit. A nucleotide sequence was analyzed using oligonucleotides consisting of nucleotide sequences represented by SEQ ID NOs: 29 and 54 as a primer and employing the purified DNA as a template according to the method described in Example 11 (1). As a result, a nucleotide sequence represented by base numbers 2241 to 3603 of the nucleotide sequence represented by SEQ ID NO: 56 was read.

(2) Analysis of full length Mycospharella tritici StOS-1 gene

[0142] A DNA having a nucleotide sequence extending toward to 5' upstream region of abase number 2241 of the nucleotide sequence represented by SEQ ID NO: 56 was cloned using SMART RACE cDNA Amplification Kit (CLONTECH) according to the instruction attached to the kit. A reaction solution was prepared by mixing 1.0  $\mu$ L of CDS-primer and 1.0  $\mu$ L of SMART IIA Oligo attached to the kit into 3  $\mu$ L (230ng) of total RNA prepared in Example 10 (2), a temperature was maintained at 70°C for 2 minutes, and maintained on ice for 2 minutes. To the reaction solution were added 2  $\mu$ L of 5×First-Strand buffer attached to the kit, 1  $\mu$ L of 20 mM DTT, 1  $\mu$ L of 10 mM dNTP Mix and 1  $\mu$ L of PowerScript Reverse Transcriptase, to mix them, and the mixture was maintained at 42°C for 1.5 hours. To the reaction solution after temperature maintenance was added 100  $\mu$ L of Tricine-EDTA buffer attached to the kit, a temperature was maintained at 72°C for 7 minutes, thus 5' RACE ready cDNA was prepared. PCR amplifying 5' upstream

region was performed using this 5' RACE ready cDNA as a template and using KOD-plus-(TOYOBO). The PCR reaction solution was prepared by mixing 2.5  $\mu$ L of 5' RACE ready cDNA, 5.0  $\mu$ L of 10×buffer, 5.0  $\mu$ L of 2 mM dNTPs, 2.0  $\mu$ L of 25 mM MgSO<sub>4</sub> and 1.0  $\mu$ L of KOD-Plus, adding 5.0  $\mu$ L of 10× Universal Primer A Mix attached to the kit and 1.0  $\mu$ L of a 10  $\mu$ M solution of an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 43 as primers, and adding sterilized distilled water to a total amount of 50  $\mu$ L. This reaction solution was maintained at 94°C for 2 minutes, and 35 cycles of incubation were repeated, each cycle comprising maintaining a temperature at 94°C for 15 seconds, then, at 68°C for 5 minutes. The amplified DNA was purified from the PCR reaction solution using QIAquick PCR Purification Kit (QIAGEN) according to the instruction attached to the kit, and then, 3'A addition was performed on the DNA according to the method described in Example 11(1). The 3'A-added DNA and the pCR2,1-TOPO cloning vector (Invitrogen) were ligated according to the instruction attached to the cloning vector, after that, which was introduced into Escherichia coli JM109 (TaKaRa). A plasmid DNA was purified from the resulting Escherichia coli transformant using QIAprep Spin Miniprep Kit (QIAGEN). A nucleotide sequence was analyzed using the resulting plasmid DNA as a template and using primers consisting of nucleotide sequences represented by SEQ ID NOs: 29, 54, and 59 to 61 according to the method described in Example 11 (1). As a result, a nucleotide sequence represented by base numbers 1 to 2240 of the nucleotide sequence represented by SEQ ID NO: 56 was read.

[0143] Further, a DNA having a nucleotide sequence extending toward to the 3' downstream region from nucleotide number 3603 of the nucleotide sequence represented by SEQ ID NO: 56 was cloned. 1. 0 μL of CDS-primer attached to the kit and 1.0  $\mu$ L of sterilized distilled water were mixed into 3  $\mu$ L (230ng) of the total RNA prepared in Example 10 (2), the mixture was maintained at 70°C for 2 minutes, and maintained on ice for 2 minutes. 3' RACE ready cDNA was prepared using the reaction solution as in preparation of 5' RACE ready cDNA. PCR amplifying 3' downstream region was performed using this 3' RACE ready cDNA as a template. A PCR reaction solution was prepared by mixing 5.0  $\mu$ L of 10imes Advantage 2 buffer attached to the kit, 1.0  $\mu$ L of 10 mM dNTP Mix and 1.0  $\mu$ L of 50imes Advantage 2 Polymerase Mix into 2.5 μL of 3' RACE ready cDNA, adding 5.0 μL of 10×Universal Primer A Mix attached to the kit as a primer, and 1. 0  $\mu L$  of a 10  $\mu M$  solution of an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 58, and adding sterilized distilled water to a total amount of 50 µL. This reaction solution was subjected to repetition of 5 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 5 seconds, then, at 72°C for 4 minutes, further repetition of 5 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 5 seconds, then, at 70°C for 10 seconds, then, at 72°C for 4 minutes, further repetition of 25 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 5 seconds, then, at 68°C for 10 seconds, then, at 72°C for 4 minutes, followed by maintaining a temperature at 72°C for 7 minutes. The PCR reaction solution and the pCR2.1-TOPO cloning vector (Invitrogen) were ligated to the vector according to the instruction attached to the kit, after that, which was introduced into Escherichia coli JM109 (TaKaRa). Aplasmid DNAwas purified from the resulting Escherichia coli transformant using QIAprep Spin Miniprep Kit (QIAGEN). A nucleotide sequence was analyzed using the resulting plasmid DNA as a template, and using a primer consisting of the nucleotide sequence represented by any of SEQ ID NOs: 29 and 54, according to the method described in Example 11 (1). As a result, a nucleotide sequence -represented by nucleotide numbers 3604 to 3924 of the nucleotide sequence represented by SEQ ID NO: 56 was read. [0144] All analyzed nucleotide sequences were joined and, as a result, the nucleotide sequence represented by SEQ ID NO: 56 was obtained. The nucleotide sequence represented by SEQ ID NO: 56 consists of 3924 bases (including termination codon), and was a nucleotide sequence encoding 1307 amino acid residues (SEQ ID NO: 55). A molecular weight of a protein having the amino acid sequence represented by SEQ ID NO: 55 was calculated to be 143276 Da.

(3) Isolation of full length Mycospharella tritici StOS-1 gene

[0145] A DNA having a nucleotide sequence encoding an amino acid sequence of Mycospharella tritici StOS-1 (hereinafter, referred to as StOS-1 DNA in some cases) was amplified by PCR using the 5' RACE ready cDNA prepared in Example 14 (2) as a template. By performing a PCR using, as a primer, an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 64 and an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 65, a DNA having the nucleotide sequence represented by SEQ ID NO: 56 was amplified, according to the method described in Example 11 (3). After the reaction, a part of the PCR reaction solution was separated by 1% agarose gel electrophoresis, and stained with ethidium bromide. It was confirmed that about 4 kb of the DNA (StOS-1 DNA) was amplified.

Example 15

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'Construction of expression plasmid of Mycospharella tritici StOS-1 gene and preparation of transformed budding yeast

[0146] The StOS-1 DNA was cloned into a pCR2 .1-TOPO cloning vector (Invitrogen). About 4 kb of the DNA (StOS-1 DNA) was purified from the PCR reaction solution prepared in Example 14 (3) using QIAquick PCR Purification Kit

(QIAGEN) according to the instruction attached to the kit. 3'A addition was performed on about 4 kb of the purified DNA (StOS-1 DNA) according to the method described in Example 11 (1). The 3'A-added about 4 kb DNA (StOS-1 DNA) and the pCR2.1-TOPO cloning vector (Invitrogen) were ligated according to the manual attached to the cloning vector to construct a plasmid pCRStOS1. A nucleotide sequence of the resulting plasmid was analyzed according to the method described in Example 11(1). As a primer, an oligonucleotide consisting of the nucleotide sequence represented by any of SEQ ID NOs: 29, 54, and 58 to 63 was used. As a result, the nucleotide sequence represented by SEQ ID NO: 56 was obtained, and it was confirmed that the plasmid pCRStOS1 was a plasmid containing the StOS-1 DNA.

[0147] The StOS-1 DNA contained in the thus prepared plasmid pCRStOS1 was cloned into a shuttle vector p415ADH replicable in yeastand Escherichia coli to construct an expression plasmid. The plasmid pCRStOS1 was digested with restriction enzymes Spel and HindIII and, on the other hand, the shuttle vector p415ADH was also digested with restriction enzymes Spel and HindIII. Each of them was separated by 0.8% agarose gel electrophoresis, a part of the gel containing the StOS-1 DNA digested with Spel and HindIII and the shuttle vector p415ADH digested with Spel and HindIII was excised, and the StOS-1 DNA and the shuttle vector were recovered from the gel using QIAquick Gel Extraction Kit (QIAGEN) according to the attached manual. The StOS-1 DNA was inserted between Spel site and HindIII site in the multicloning site of the shuttle vector using Ligation Kit Ver.2 (TaKaRa) according to the manual attached to the kit, whereby, an expression plasmid pADHStOS1 was constructed. A nucleotide sequence of the resulting expression plasmid was analyzed according to the method described in Example 11 (1). As a primer, an oligonucleotide consisting of the nucleotide sequence represented by any of SEQ ID NO: 58 to 65 was used. As a result, the nucleotide sequence represented by SEQ ID NO: 56 was obtained, and it was confirmed that the expression plasmid pADHStOS1 harbored a DNA having a nucleotide sequence encoding an amino acid sequence of StOS-1. [0148] The prepared expression plasmid pADHStOS1 was introduced into budding yeast AH22 strain and TM182 strain according to the method described in Example 2. By utilizing the disappearance of leucine auxotrophy in the resulting transformed budding yeast, the transformed budding yeast AH22 strain (AH22-StOS1) was selected on a Glu-Leu agar medium, and the transformed budding yeast TM182 strain (TM182-StOS1) was selected on a Gal-Ura-Leu agar medium. It was confirmed that the resulting TM182-StOS1 grows even when transplanted to a Glu-Ura-Leu medium.

#### Example 16

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Antifungal compound sensitivity test of transformed budding yeast TM182-StOS1

[0149] The transformed budding yeast AH22-StOS1 prepared in Example 15 was cultured while shaking at 30°C for 18 hours in a Glu-Leu medium. As a control, the AH22 strain was similarly cultured while shaking at 30°C for 18 hours in a Glu medium. The absorbance at 600 nm of each grown transformed budding yeast in a cell suspension was measured, and cell suspension diluted with sterilized distilled water to the absorbance of 0.1 was prepared. Further, a cell suspension in which the aforementioned cell suspension of the transformed budding yeast AH22-StOS1 was diluted 50-fold with a Glu-Leu medium, and a cell suspension in which the aforementioned cell suspension of the AH22 strain was diluted 50-fold with a Glu medium were prepared.

[0150] A solution in which each of Compounds (1) to (3) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 6 ppm, a solution in which each of Compounds (4) and (5) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 2000 ppm, and a solution in which Compounds (6) and (7) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 20 ppm were prepared, and two microplates were prepared in which each 1.0 µL per well of the Compound solution and DMSO as a control were dispensed. In one microplate among them, each 100 µL of cell suspensions of the transformed budding yeast AH22-StOS1 which had been prepared by dilution as described above was dispensed, and cultured by allowing to stand at 30°C for 28 hours. In another microplate, each 100 µL of cell suspensions of the control yeast AH22 strain which had been prepared by dilution as described above was dispensed, and cultured by allowing to stand at 30°C for 24.5 hours. After culturing, the absorbance at 600 nm of each well was measured with a microplate reader.

[0151] Similarly, the transformed budding yeast TM182-StOS1 prepared in Example 15 was cultured at 30°C for 18 hours in a Glu-Ura-Leu medium. The absorbance at 600 nm of the grown transformed budding yeast in a cell suspension was measured, and a cell suspension diluted with sterilized distilled water to the absorbance of 0.1 was prepared. Further, a cell suspension in which the transformed budding yeast TM182-StOS1 was diluted 50-fold with a Glu-Ura-Leu medium and, as a control, a cell suspension in which the yeast was diluted 50-fold with a Gal-Ura-Leu medium were prepared.

[0152] A solution in which each of Compounds (1) to (3) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 6 ppm, a solution in which each of Compounds (4) and (5) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 2000 ppm, and a solution in which each of Compounds (6) and (7) was dissolved in dimethylsulfoxide

(DMSO) to the concentration of 20 ppm were prepared, and two microplates were prepared in which each  $2.0~\mu$ L per well of the Compound-DMSO solution and DMSO as a control were dispensed into 2 wells. In one microplate among them, each  $100~\mu$ L of cell suspensions of the transformed budding yeast TM182-StOS1 which had been prepared by dilution with a Glu-Ura-Leu medium as described above was dispensed, and cultured by allowing to stand at  $30^{\circ}$ C for  $26.5~\mu$ L in another microplate, as described above, as a control, each  $100~\mu$ L of cell suspensions of the transformed budding yeast TM182-StOS1 which had been prepared by dilution with a Gal-Ura-Leu medium was dispensed, and cultured at  $30^{\circ}$ C for  $49.5~\mu$ C for 49.5~

[0153] A degree of growth of each transformed budding yeast cultured in the presence of Compounds (1) to (7) is shown in Table 6. A degree of growth of the transformed budding yeast is expressed as a relative value in percentage, letting the absorbance at 600 nm at the concentration of the Compound of 0 ppm to be 100. It was confirmed that an inhibiting degree of growth of the transformed budding yeast TM182-StOS1 by each test substance was grater than an inhibiting degree of growth of the transformed budding yeast AH22-StOS1 by each test substance, and the transformed budding yeast TM182-StOS1 was a transformed cell with the enhanced sensitivity to an antifungal compound as compared with the transformed budding yeast AH22-StOS1.

Table 6

Degree of growth of budding yeast					
	AH22	AH22-St OS1 TM182-StOS1			
Test substance (final concentration)	Glu medium	Glu-Leu medium	Gal-Ura-Leu medium	Gul-Ura-Leu mediun	
Compoud (1) (0.6 ppm)	99	101	101	67	
Compoud (2) (0.6 ppm)	94	100	97	23	
Compoud (3) (0.6 ppm)	96	98	94	19	
Compoud (4) (20 ppm)	96	91	99	7	
Compoud (5) (20 ppm)	80	76	74	6	
Compoud (6) (0.2 ppm)	92	93	97	6	
Compoud (7) (0.2 ppm)	91	91	91	9	

Example 17

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Isolation of Thanatephorus cucumeris RsOS-1 gene

(1) Analysis of Thanatephorus cucumeris RsOS-1 gene fragment

[0154] The amplified DNA was purified from the reaction solution of PCR which had been performed using a primer pair 3 and using a cDNA of Thanatephorus cucumeris as a template in Example 10 (5), using QlAquick PCR Purification Kit (QlAGEN) according to the instruction attached to the kit. 3'A addition was performed on the purified DNA according to the method described in Example 11 (1). The 3' A-added DNA and the pCR2.1-TOPO cloning vector (Invitrogen) were ligated according to the instruction attached to the cloning vector, and this was introduced into Escherichia coli JM109 (TaKaRa).

[0155] DNA was purified from the resulting Escherichia coli transformant by colony PCR using Ex Taq HS (TaKaRa). The PCR reaction solution (15  $\mu$ L) was prepared by mixing 1.5  $\mu$ L of 10×buffer, 2.25  $\mu$ L of 10 mM dNTPs, 0.15  $\mu$ L of Ex Taq HS, each 0.4  $\mu$ L of a 10  $\mu$ M solution of an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 28 and a 10  $\mu$ M solution of an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 29, and 10.3  $\mu$ L of sterilized distilled water, and adding a part of the Escherichia coli transformant colony thereto. PCR was performed under the amplifying conditions that this reaction solution was maintained at 97°C for 2 minutes and, thereafter, 35 cycles of incubation were repeated, each cycle comprising maintaining a temperature at 97°C for 15 seconds, then, at 55°C for 15 seconds, then, at 72°C for 3 minutes. The amplified DNA was purified from the PCR reaction solution after temperature maintenance using QIAquick PCR Purification Kit (QIAGEN) according to the instruction attached to the kit. A nucleotide sequence was analyzed using oligonucleotides consisting of nucleotide sequences represented by SEQ ID NOs: 28 and 29 as a primer and employing the purified DNA as a template according to the method described in Example 11 (1). As a result, a nucleotide sequence represented by base numbers 2838 to

3165 of the nucleotide sequence represented by SEQ ID NO: 69 was read.

(2) Analysis of full length Thanatephorus cucumeris RsOS-1 gene

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[0156] A DNA having a nucleotide sequence extending toward to 3' downstream region of a base number 3165 of the nucleotide sequence represented by SEQ ID NO: 69 was cloned using SMART RACE cDNA Amplification Kit (CLONTECH) according to the instruction attached to the kit. A reaction solution was prepared by mixing 1.0 μL of CDS-primer and 1.0  $\mu$ L of sterilized distilled water attached to the kit into 3  $\mu$ L (253ng) of total RNA prepared in Example 10 (3), a temperature was maintained at 70°C for 2 minutes, and maintained on ice for 2 minutes. To the reaction solution were added 2  $\mu$ L of 5×First-Strand buffer attached to the kit, 1  $\mu$ L of 20 mM DTT, 1  $\mu$ L of 10 mM dNTP Mix and 1 μL of PowerScript Reverse Transcriptase, to mix them, and the mixture was maintained at 42°C for 1. 5 hours. To the reaction solution after temperature maintenance was added 100 µL of Tricine-EDTA buffer attached to the kit. a temperature was maintained at 72°C for 7 minutes, thus 3' RACE ready cDNA was prepared. PCR amplifying 3' downstream region was performed using this 3' RACE ready cDNA as a template and using KOD-plus-(TOYOBO). The PCR reaction solution was prepared by mixing 2.5  $\mu$ L of 3' RACE ready cDNA, 5.0  $\mu$ L of 10×buffer, 5.0  $\mu$ L of 2 mM dNTPs, 2.0  $\mu$ L of 25 mM MgSO<sub>4</sub> and 1.0  $\mu$ L of KOD-Plus, adding 5.0  $\mu$ L of 10 $\times$  Universal Primer A Mix attached to the kit and 1.0  $\mu$ L of a 10  $\mu$ M solution of an oligonucleotide consisting of the nucleotide sequence represented by SEQ IDNO: 70 as primers, and adding sterilized distilled water to a total amount of 50 μL. This reaction solution was maintained at 94°C for 2 minutes, and 35 cycles of incubation were repeated, each cycle comprising maintaining a temperature at 94°C for 15 seconds, then, at 68°C for 6 minutes. The amplified DNA was purified from the PCR reaction solution using QIAquick PCR Purification Kit (QIAGEN) according to the instruction attached to the kit, and then, 3' A addition was performed on the DNA according to the method described in Example 11(1). The 3'A-added DNA and the pCR2.1-TOPO cloning vector (Invitrogen) were ligated according to the instruction attached to the cloning vector, after that, which was introduced into Escherichia coli JM109 (TaKaRa). A plasmid DNA was purified from the resulting Escherichia coli transformant using QIAprep Spin Miniprep Kit (QIAGEN). A nucleotide sequence was analyzed using the resulting plasmid DNA as a template and using primers consisting of nucleotide sequences represented by SEQ IDNOs: 28, 29, and 73 to 76 according to the method described in Example 11(1). As a result, a nucleotide sequence represented by base numbers 3119 to 4317 of the nucleotide sequence represented by SEQ ID NO: 69 was read. [0157] Further, a DNA having a nucleotide sequence extending toward to the 5' upstream region from nucleotide number 2838 of the nucleotide sequence represented by SEQ IDNO: 69 was cloned. 1.0 μL of CDS-primer attached to the kit and 1.0 μL of SMART IIA Oligo were mixed into 3 μL (253ng) of the total RNA prepared in Example 10 (3), the mixture was maintained at 70°C for 2 minutes, and maintained on ice for 2 minutes. 5' RACE ready cDNA was prepared using the reaction solution as in preparation of 3' RACE ready cDNA. PCR amplifying 5' upstream region was performed using this 5' RACE ready cDNA as a template and using KOD-plus-(TOYOBO). The PCR reaction solution was prepared by mixing 2.5 μL of 5' RACE ready cDNA, 5.0 μL of 10×buffer, 5.0 μL of 2 mM dNTPs, 2.0 μL -of 25 mM MgSO<sub>4</sub> and 1.0  $\mu$ L of KOD-Plus, adding 5. 0  $\mu$ L of 10 $\times$ Universal Primer A Mix attached to the kit and 1.0 μL of a 10 μM solution of an oligonucleotide consisting of the nucleotide sequence represented by SEQ IDNO: 71 as primers, and adding sterilized distilled water to a total amount of 50 µL. This reaction solution was maintained at 94°C for 2 minutes, and 35 cycles of incubation were repeated, each cycle comprising maintaining a temperature at 94°C for 15 seconds, then, at 68°C for 6 minutes. Using the resulting PCR reaction solution as a template, the PCR reaction solution for a futher PCR was prepared by adding 5.0  $\mu$ L of 10 $\times$ buffer, 5.0  $\mu$ L of 2 mM dNTPs, 2.0  $\mu$ L of 25 mM MgSO<sub>4</sub> and 1.0 μL of KOD-Plus, 1.0 μL of 10 μM Nested universal primer attached to the kit and 1.0 μL of a 10 μM solution of an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 72 as primers, and adding sterilized distilled water to a total amount of 50 µL. This reaction solution was maintained at 94°C for 2 minutes, and 20 cycles of incubation were repeated, each cycle comprising maintaining a temperature at 94°C for 15 seconds, then, at 68°C for 6 minutes. The PCR reaction solution and the pCR2.1-TOPO cloning vector (Invitrogen) were ligated to the vector according to the instruction attached to the kit, after that, which was introduced into Escherichia coli JM109 (TaKaRa). A plasmid DNA was purified from the resulting Escherichia coli transformant using QIAprep Spin Miniprep Kit (QIAGEN). A nucleotide sequence was analyzed using the resulting plasmid DNA as a template, and using a primer consisting of the nucleotide sequence represented by any of SEQ ID NOs: 28, 29, and 77 to 82, according to the method described in Example 11 (1). As a result, a nucleotide sequence represented by nucleotide numbers 1 to 3042 of the nucleotide sequence represented by SEQ ID NO: 69 was read. [0158] All analyzed nucleotide sequences were joined and, as a result, the nucleotide sequence represented by SEQ

ID NO: 69 was obtained. The nucleotide sequence represented by SEQ ID NQ: 69 consists of 4317 bases (including termination codon), and was a nucleotide sequence encoding 1438 amino acid residues (SEQ ID NO: 68). A molecular weight of a protein having the amino acid sequence represented by SEQ ID NO: 68 was calculated to be 155296 Da.

(3) Isolation of full length Thanatephorus cucumeris RsOS-1 gene

[0159] A DNA having a nucleotide sequence encoding an amino acid sequence of Thanatephorus cucumeris RsOS-1 (hereinafter, referred to as RsOS-1 DNA in some cases) was amplified by PCR using a cDNA of Thanatephorus cucumeris prepared in Example 10 (5) as a template. By performing a PCR using, as a primer, an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 85 and an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 86, a DNA having the nucleotide sequence represented by SEQ ID NO: 69 was amplified, according to the method described in Example 11 (3). After the reaction, a part of the PCR reaction solution was separated by 1% agarose gel electrophoresis, and stained with ethidium bromide. It was confirmed that about 4 kb of the DNA (RsOS-1 DNA) was amplified.

#### Example 18

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Construction of expression plasmid of Thanatephorus cucumeris RsOS-1 gene and preparation of transformed budding yeast

[0160] The RsOS-1 DNA was cloned into a pCR2. 1-TOPO cloning vector (Invitrogen). About 4 kb of the DNA (RsOS-1 DNA) was purified from the PCR reaction solution prepared in Example 17 (3) using QIAquick PCR Purification Kit (QIAGEN) according to the instruction attached to the kit. 3'A addition was performed on about 4 kb of the purified DNA (StOS-1 DNA) according to the method described in Example 11 (1). The 3'A-added about 4 kb DNA (RsOS-1 DNA) and the pCR2. 1-TOPO cloning vector (Invitrogen) were ligated according to the manual attached to the cloning vector to construct a plasmid pCRRsOS1. A nucleotide sequence of the resulting plasmid was analyzed according to the method described in Example 11(1). As a primer, an oligonucleotide consisting of the nucleotide sequence represented by any of SEQ ID NOs: 28, 29, 70 to 73, 75, 77, 78, and 81 to 84 was used. As a result, the nucleotide sequence represented by SEQ ID NO: 69 was obtained, and it was confirmed that the plasmid pCRRsOS1 was a plasmid containing the RsOS-1 DNA.

[0161] The RsOS-1 DNA contained in the thus prepared plasmid pCRRsOS1 was cloned into a shuttle vector p415ADH replicable in yeastand Escherichia coli to construct an expression plasmid. The plasmid pCRRsOS1 was digested with restriction enzymes Spel and HindIII and, on the other hand, the shuttle vector p4J.5ADH was also digested with restriction enzymes Spel and HindIII. Each of them was separated by 0.8% agarose gel electrophoresis, a part of the gel containing the RsOS-1 DNA digested with Spel and HindIII and the shuttle vector p415ADH digested with Spel and HindIII was excised, and the RsOS-1 DNA and the shuttle vector were recovered from the gel using QIAquick Gel Extraction Kit (QIAGEN) according to the attached manual. The RsOS-1 DNA was inserted between Spel site and HindIII site in the multicloning site of the shuttle vector using Ligation Kit Ver.2 (TaKaRa) according to the manual attached to the kit, whereby, an expression plasmid pADHRsOS1 was constructed. A nucleotide sequence of the resulting expression plasmid was analyzed according to the method described in Example 11 (1). As a primer, an oligonucleotide consisting of the nucleotide sequence represented by any of SEQ ID NO: 70 to 73, 75, 77, 78, 81 to 84, 87 and 88 was used. As a result, the nucleotide sequence represented by SEQ ID NO: 69 was obtained, and it was confirmed that the expression plasmid pADHRsOS1 harbored a DNA having a nucleotide sequence encoding an amino acid sequence of RsOS-1.

[0162] The prepared expression plasmid pADHRsOS1 was introduced into budding yeast AH22 strain and TM182 strain according to the method described in Example 2. By utilizing the disappearance of leucine auxotrophy in the resulting transformed budding yeast, the transformed budding yeast AH22 strain (AH22-RsOS1) was selected on a Glu-Leu agar medium, and the transformed budding yeast TM182 strain (TM182-RsOS1) was selected on a Gal-Ura-Leu agar medium. It was confirmed that the resulting TM182-RsOS1 grows even when transplanted to a Glu-Ura-Leu medium.

#### Example 19

Antifungal compound sensitivity test of transformed budding yeast TM182-RsOS1

[0163] The transformed budding yeast AH22-RsOS1 prepared in Example 18 was cultured while shaking at 30°C for 18 hours in a Glu-Leu medium. As a control, the AH22 strain was similarly cultured while shaking at 30°C for 18 hours in a Glu medium. The absorbance at 600 nm of each grown transformed budding yeast in a cell suspension was measured, and cell suspension diluted with sterilized distilled water to the absorbance of 0.1 was prepared. Further, a cell suspension in which the aforementioned cell suspension of the transformed budding yeast AH22-RsOS1 was diluted 50-fold with a Glu-Leu medium, and a cell suspension in which the aforementioned cell suspension of the AH22 strain was diluted 50-fold with a Glu medium were prepared.

[0164] A solution in which each of Compounds (1) to (5) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 600ppm, and a solution in which Compounds (6) and (7) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 60 ppm were prepared, and two microplates were prepared in which each 1.0  $\mu$ L per well of the Compound solution and DMSO as a control were dispensed. In one microplate among them, each 100  $\mu$ L of cell suspensions of the transformed budding yeast AH22-RsOS1 which had been prepared by dilution as described above was dispensed, and cultured by allowing to stand at 30 °C for 29.8 hours. In another microplate, each 100  $\mu$ L of cell suspensions of the control yeast AH22 strain which had been prepared by dilution as described above was dispensed, and cultured by allowing to stand at 30 °C for 24.8 hours. After culturing, the absorbance at 600 nm of each well was measured with a microplate reader.

[0165] Similarly, the transformed budding yeast TM182-RsOS1 prepared in Example 18 was cultured at 30°C for 18 hours in a Glu-Ura-Leu medium. The absorbance at 600 nm of the grown transformed budding yeast in a cell suspension was measured, and a cell suspension diluted with sterilized distilled water to the absorbance of 0.1 was prepared. Further, a cell suspension in which the transformed budding yeast TM182-RsOS1 was diluted 50-fold with a Glu-Ura-Leu medium. As a control, the transformed budding yeast TM182-RsOS1 was cultured at 30°C for 18 hours in a Gal-Ura-Leu medium. The absorbance at 600 nm of the grown transformed budding yeast in a cell suspension was measured, and a cell suspension diluted with sterilized distilled water to the absorbance of 0.1 was prepared. Further, a cell suspension in which the transformed budding yeast TM182-RsOS1 was diluted 50-fold with a Gal-Ura-Leu medium. [0166] A solution in which each of Compounds (1) to (5) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 600 ppm, and a solution in which each of Compounds (6) and (7) was dissolved in dimethylsulfoxide (DMSO) to the concentration of 60 ppm were prepared, and two microplates were prepared in which each 2.0 µL per well of the Compound-DMSO solution and DMSO as a control were dispensed into 2 wells. In one microplate among them, each 100 µL of cell suspensions of the transformed budding yeast TM182-RsOS1 which had been prepared by dilution with a Glu-Ura-Leu medium as described above was dispensed, and cultured by allowing to stand at 30°C for 26.8 hours. In another microplate, as described above, as a control, each 100  $\mu$ L of cell suspensions of the transformed budding yeast TM182-RsOS1 which had been prepared by dilution with a Gal-Ura-Leu medium was dispensed, and cultured at 30°C for 42.5 hours. After culturing, the absorbance at 600 nm of each well was measured with a microplate reader.

[0167] A degree of growth of each transformed budding yeast cultured in the presence of Compounds (1) to (7) is shown in Table 7. A degree of growth of the transformed budding yeast is expressed as a relative value in percentage, letting the absorbance at 600 nm at the concentration of the Compound of 0 ppm to be 100. It was confirmed that an inhibiting degree of growth of the transformed budding yeast TM182-RsOS1 by each test substance was grater than an inhibiting degree of growth of the transformed budding yeast AH22-RsOS1 by each test substance, and the transformed budding yeast TM182-RsOS1 was a transformed cell with the enhanced sensitivity to an antifungal compound as compared with the transformed budding yeast AH22-RsOS1.

Table 7

	Degree of gro	owth of budding year	st					
	AH22	AH22-Rs	TM182-RsOS1					
Test substance (final concentration)	Glu medium	Glu-Leu medium	Gal-Ura-Leu medium	Gul-Ura-Leu medium				
Compoud (1) (6.0 ppm)	88	. 103	108	15				
Compoud (2) (6.0 ppm)	92	101	96	11				
Compoud (3) (6.0 ppm)	82	101	101	27				
Compoud (4) (6.0 ppm)	83	89	88	17				
Compoud (5) (6.0 ppm)	78	85	101	9				
Compoud (6) (0.6 ppm)	79	79	100	12				
Compoud (7) (0.6 ppm)	85	101	99	31				

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#### Example 20

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Isolation of a gene of the present histidine kinase of Phytophthora infestans (hereinafter, referred to PiOS-1 gene)

(1) Analysis of Phytophthora infestans PiOS-1 gene fragment

[0168] The amplified DNA was purified from the reaction solution of PCR which had been performed using a primer pair 6 and using a cDNA of Phytophthora infestans as a template in Example 10 (5), using QIAquick PCR Purification Kit (QIAGEN) according to the instruction attached to the kit. 3'A addition was performed on the purified DNA according to the method described in Example 11 (1). The 3'A added DNA and the pCR2.1-TOPO cloning vector (Invitrogen) were ligated according to the instruction attached to the cloning vector, after that, which was introduced into Escherichia coli JM109 (TaKaRa).

[0169] A DNA was amplified from the resulting Escherichia coli transformant by colony PCR using Ex Taq HS (TaKa-Ra). The PCR reaction solution (15  $\mu$ L) was prepared by mixing 1.5  $\mu$ L of 10×buffer, 2.25  $\mu$ L of 10 mM dNTPs, 0.15  $\mu$ L of Ex Taq HS, each 0.4  $\mu$ L of a 10  $\mu$ M solution of an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 28 and a 10  $\mu$ M solution of an oligonucleotide consisting of the nucleotide sequence represented by SEQ ID NO: 29, and 10. 3  $\mu$ L of sterilized distilled water, and adding a part of the Escherichia coli transformant colony thereto. PCR was performed under the amplifying conditions that this reaction solution was maintained at 97°C for 2 minutes, and 35 cycles of incubation were repeated, each cycle comprising maintaining a temperature at 97°C for 15 seconds, then, at 55°C for 15 seconds, further, at 72°C for 3 minutes. The amplified DNA was purified from the PCR reaction solution after temperature maintenance using QIAquick PCR purification Kit (QIAGEN) according to the manual attached to the kit. A nucleotide sequence was analyzed using the purified DNA as a template and using oligonucleotides consisting of the nucleotide sequence represented by any of SEQ ID NOs: 28 and 29 as primers according to the method described in Example 11 (1). As a result, a nucleotide sequence represented by SEQ ID NO: 89 containing a nucleotide sequence of an oligonucleotide used as a primer pair 6 was read.

(2) Analysis of full length Phytophthora infestans PiOS-1 gene

[0170] A DNA having a nucleotide sequence extending toward to 5' upstream region of a nucleotide sequence represented by SEQ ID NO: 89 is cloned using SMART RACE cDNA Amplification Kit (CLONTECH) according to the instruction attached to the kit. A reaction solution is prepared by mixing 1.0 µL of CDS-primer attached to the kit and  $1.0\,\mu L$  of SMART IIA Oligo into  $3\,\mu L$  (200ng) of the total RNA prepared in Example 10 (4), a temperature is maintained at 70°C for 2 minutes, and is maintained on ice for 2 minutes. To the reaction solution are added 2 µL of 5×First-Strand buffer attached to the kit, 1 µL of 20 mM DTT, 1 µL of 10 mM dNTP Mix and 1 µL of PowerScript Reverse Transcriptase to mix them, and the mixture is maintained at 42°C for 1.5 hours. To the reaction solution after temperature maintenance -is added 100  $\mu$ L of Tricine-EDTA buffer attached to the kit, a temperature is maintained at 72°C for 7 minutes, and 5' RACE ready cDNA is prepared. PCR amplifying 5' upstream region is performed using this 5' RACE ready cDNAas a template and using KOD-plus- (TOYOBO). The PCR reaction solution is prepared by mixing 2.5 μL of 5' RACE ready cDNA, 5.0 μL of 10×buffer, 5.0 μL of 2 mM dNTPs, 2.0 μL of 25 mM MgSO<sub>4</sub> and 1.0 μL of KOD-Plus-, adding 5.0 μL of 10×Universal primer A Mix attached to the kit as a primer and 1.0 μL of a 10 μM solution of an oligonucleotide consisting of 20 to 30 bases selected from complementary sequences of the nucleotide sequence represented by SEQ ID NO: 89, and adding sterilized distilled water to a total amount of 50 μL. This reaction solution is maintained at 94°C for 2 minutes, and further 35 cycles of incubation are repeated, each cycle comprising maintaining a temperature at 94°C for 15 seconds, then, at 68°C for 5 minutes. The amplified DNA is purified from the PCR reaction solution using QIAquick PCR Purification Kit (QIAGEN) according to the instruction attached to the kit, and 3' A addition is performed on the DNA according to the method described in Example 11 (1). 3' A added DNA and the pCR2 . 1-TOPO cloning vector (Invitrogen) are ligated according to the instruction attached to the cloning vector, after that, which is introduced into Escherichia coli JM109 (TaKaRa). A plasmid DNA is purified from the resulting Escherichia coli transformant using QIAprep Spin Miniprep Kit (QIAGEN). A nucleotide sequence is analyzed using the resulting plasmid DNA as a template and using primers consisting of the nucleotide sequence represented by any of SEQ ID NOs: 28, 29, and the like according to the method described in Example 11(1). As a result, a nucleotide sequence of the 5'-terminal region including a translation initiation codon of an os-1 homologous gene of Phytophthora infestans, that is, gene of Phytophthora infestans encoding osmosensing histidine kinase having no transmembrane region (PiOS1) can be read. [0171] Further, a DNA having a nucleotide sequence extending to 3' downstream region of the nucleotide sequence represented by SEQ ID NO: 89 is cloned. 1.0 μL of CDS-primer attached to the kit and 1.0 μL of sterilized distilled water are mixed into 3 μL (200ng) of the total RNA prepared in Example 10 (4), a temperature is maintained at 70°C for 2 minutes, and is maintained on ice for 2 minutes. 3' RACE ready cDNA is prepared using the reaction solution as in preparation of 5' RACE ready cDNA. PCR amplifying 3' downstream region is performed using this 3' RACE ready

cDNA as a template. The PCR reaction solution is prepared by mixing 5.0 µL of 10×Advantage 2 buffer attached to the kit, 1.0  $\mu$ L of 10 mM dNTP Mix and 1.0  $\mu$ L of 50×Advantage 2 polymerase Mix into 2.5  $\mu$ L of 5' RACE ready cDNA, adding 5. 0 μL of 10×Universal Primer A Mix attached to the kit, and 1.0 μL of a 10 μM solution of an oligonucleotide consisting of 20 to 30 bases selected from the nucleotide sequence represented by SEQ ID NO: 89 as primers, and adding sterilized distilled water to a total amount of 50 µL. This reaction solution is subjected to repetition of 5 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 5 seconds, then, at 72°C for 4 minutes, further repetition of 5 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 5 seconds, then, at 70°C for 10 seconds then, at 72°C for 4 minutes, further repetition of 25 cycles of incubation, each cycle comprising maintaining a temperature at 94°C for 5 seconds, then, at 68°C for 10 seconds, then, at 72°C for 4 minutes, followedbymaintaining a temperature at 72°C for 7 minutes. The PCR reaction solution and the pCR2.1-TOPO cloning vector (Invitrogen) are ligated according to the instruction attached to the cloning vector, after that, which is introduced into Escherichia coli JM109 (TaKaRa). A plasmid DNA is purified from the resulting Escherichia coli transformant using QIAprep Spin Miniprep Kit (QIAGEN). A nucleotide sequence is analyzed using the resulting plasmid DNA as a template and using primers consisting of the nucleotide sequence represented by any of SEQ ID NOs: 28, 29, and the like according to the method described in Example 11 (1). As a result, a nucleotide sequence of the 3'-terminal region including a translation termination codon of a Phytophthora infestans PiOS1 gene is read.

[0172] By ligating all analyzed nucleotide sequences, full nucleotide sequence of Phytophthora infestans PiOS-1 gene including nucleotide sequence represented by SEQ ID NO: 89 is confirmed.

(3) Isolation of full length Phytophthora infestans PiOS1 gene

[0173] A DNA having a nucleotide sequence encoding an amino acid sequence of Phytophthora infestans PiOS1 (hereinafter, referred to as PiOS-1 DNA) is amplified by PCR using the cDNA prepared in Example 10 (4) as a template. Using as primers an oligonucleotide comprising a nucleotide sequence in which a nucleotide sequence ACGACAGT is added to the 5'-terminal end of a nucleotide sequence from the 5'-terminal end to the 20th base including the initiation codon of a nucleotide sequence of Phytophthora infestans PiOS-1 gene obtained in Example 20 (2), and an oligonucleotide having a nucleotide sequence complementary to a nucleotide sequence in which a nucleotide sequence AAGCTTCAG is added to the 3'-terminal end of a nucleotide sequence of from the 3'-terminal end to the 20th base including the termination codon of a nucleotide sequence of Phytophthora infestans PiOS-1 gene obtained in Example 20 (2), a PCR is performed according to the method described in Example 11 (3). DNA containing a nucleotide sequence encoding an amino acid sequence of Phytophthora infestans PiOS-1, and having a recognition sequence of a restriction enzyme Spel immediately before an initiation codon, and having a recognition sequence of a restriction enzyme HindIII immediately after a termination codon is amplified. A part of the PCR reaction solution after the reaction is separated by 1% agarose gel electrophoresis, and stained with ethidium bromide. It is confirmed that the about 4 kb PiOS-1 DNA is amplified.

Example 21

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Construction of expression plasmid of Phytophthora infestans PiOS-1 gene and preparation of transformed budding yeast

[0174] The PiOS-1 DNA is cloned into the pCR2.1-TOPO cloning vector (Invitrogen). An about 4 kb DNA (PiOS-1 DNA) is purified from the PCR reaction solution prepared in Example 20 (3) using QIAquick PCR Purification Kit (QIA-GEN) according to the manual attached to the kit. 3'A addition is performed on the about 4 kb purified DNA according to the method described in Example 11 (3). The about 4 kb 3'A-added DNA (PiOS-1 DNA) and the pCR2.1-TOPO cloning vector (Invitrogen) are ligated according to the instruction attached to the cloning vector, whereby, the plasmid pCRPiOS1 is constructed. A nucleotide sequence of the resulting plasmid is analyzed by the method described in Example 11 (1). As a primer, oligonucleotides consisting of the nucleotide sequence represented by any of SEQ ID NOs: 28 and 29 are used. As a result, it is confirmed that the plasmid pCRPiOS1 is a plasmid harboring PiOS-1 DNA containing the nucleotide sequence represented by SEQ ID NO: 89.

[0175] The Phytophthora infestans PiOS-1 gene contained in the thus prepared plasmid pCR PiOS1 is cloned into a shuttle vector p415ADH replicable in yeast and Escherichia coli, whereby, an expression plasmid is constructed. The plasmid pCRPiOS1 is digested with restriction enzymes Spel and HindIII and, on the other hand, the shuttle vector p415ADH is also digested with restriction enzymes Spel and HindIII. These are separated by 0.8% agarose gel electrophoresis, respectively, thereafter, a part of the gel containing the PiOS-1 DNA digested with restriction enzymes Spel and HindIII and the shuttle vector p415ADH digested with Spel and HindIII is excised, and the PiOS-1 DNA and the shuttle vector are recovered from the gel using QIAquick Gel Extraction Kit (QUAGEN) according to the manual attached to the kit, the PiOS-1 DNA is

inserted between Spel site and HindIII site in the multicloning site of the shuttle vector, whereby, the expression plasmid pADHPiOS1 is constructed. A nucleotide sequence of the resulting expression plasmid is analyzed according to the method described in Example 11 (1). As a primer, oligonucleotides consisting of the nucleotide sequence represented by any of SEQ ID NOs: 28 and 29 are used. As a result, it is confirmed that the expression plasmid pADHPiOS1 is a plasmid harboring the PiOS-1 DNA containing the nucleotide sequence represented by SIQ ID NO: 89.

[0176] The prepared expression plasmid pADH PiOS1 is gene-introduced into budding yeast AH22 strain and TM182 strain according to the method described in Example 2. By utilizing disappearance of leucine auxotrophy in the resulting transformed budding yeast, the transformed budding yeast AH22 strain (AH22-PiOS1) is selected on a Glu-Leu agar medium, and the transformed budding yeast TM182 strain (TN182-PiOS1) is selected on a Gal-Ura-Leu agar medium. It is confirmed that the resulting TM182-PiOS1 grows even when transplanted to a Glu-Ura-Leumedium.

#### Example 22

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Antifungal compound sensitivity test of transformed budding yeast TM182-PiOS1

[0177] The transformed budding yeast AH22-PiOS1 prepared in Example 21 is cultured while shaking at 30°C in a Glu-Leu medium. As a control, the AH22 strain is similarly cultured while shaking at 30°C in a Glu medium. The absorbance at 600 nm of each of grown transformed budding yeasts in a cell suspension is measured, and a cell suspension diluted with sterilized distilled water to the absorbance of 0.1 is prepared. Further, a cell suspension inwhich the aforementioned suspension of the transformedbudding yeast AH22-PiOS1 is diluted 50-fold with a Glu-Leu medium, and a cell suspension in which the aforementioned suspension of the AH22 strain is diluted 50-fold with a Glu medium are prepared.

[0178] A solution in which each of Compounds (1) to (7) is dissolved in dimethylsulfoxide (DMSO) is prepared, and two microplates are prepared in which each 1.0  $\mu$ L per well of each of the Compound Solution and DMSO as a control are dispensed into two wells. In one microplate among them, each 100  $\mu$ L of cell suspensions of the transformed budding yeast AH22-PiOS1 which has been prepared by dilution as described above is dispensed, and is cultured by allowing to stand at 30°C. In another microplate, each 100  $\mu$ L of cell suspensions of the control yeast AH22 strain which has been prepared by dilution as described above is dispensed, and is cultured by allowing to stand at 30°C. After culturing, the absorbance at 600 nm of each well is measured with a microplate reader.

[0179] Similarly, the transformed budding yeast TM182-PiOS1 prepared in Example 21 is cultured at 30°C in a Glu-Ura-Leu medium. The absorbance at 600 nm of a cell suspension of the grown transformed budding yeast is measured, and a cell suspension diluted with sterilized distilled water to the absorbance of 0.1 is prepared. Further, a cell suspension in which the aforementioned cell suspension of the transformed budding yeast TM182-PiOS1 is diluted 50-fold with a Glu-Ura-Leu medium and, as a control, a cell suspension in which the aforementioned cell suspension is diluted 50-fold with a Gal-Ura-Leu medium are prepared.

**[0180]** A solution in which each of Compounds (1) to (7) is dissolved in dimethylsulfoxide (DMSO) is dissolved is prepared, and two microplates are prepared in which each 1.0  $\mu$ L per well of the Compound solution and DMSO as a control are dispensed. In one microplate among them, each 100  $\mu$ L of cell suspensions of the transformed budding yeast TM182-PiOS1 which has been prepared by dilution with a Glu-Ura-Leu medium as described above is dispensed, and is cultured by allowing to stand at 30°C. In another microplate as described above, as a control, each 100  $\mu$ L of cell suspensions of the transformed budding yeast TM182-PiOS1 which has been prepared by dilution with a Gal-Ura-Leu medium is dispensed, and is cultured by allowing to stand at 30°C. After culturing, the absorbance at 600 nm of each well is measured with a microplate reader.

[0181] It is confirmed that an inhibiting degree of growth of the transformed budding yeast TM182-PiOS1 by each test substance is greater than an inhibiting degree of growth of the transformed budding yeast AH22-PiOS1 by each test substance, and the transformed budding yeast TM182-PiOS1 is a transformed cell with the enhanced sensitivity to an antifungal compound as compared with the transformed budding yeast AH2-PiOS1.

[0182] The compositions of media used in the present invention are described below.

(a) Glu-medium

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Becto-yeast nitrogen base without amino acids 6.7 g, Glucose 20 g, Drop-out mix(1) 2.0 g, Distilled water 1000 ml (b) Glu-Leu medium

Bacto-yeast nitrogen base without amino acids 6.7 g, Glucose 20 g, Drop-out mix(2) 2.0 g, Distilled water 1000 mł (c) Glu-Ura-Leu medium

Bacto-yeast nitrogen base without amino acids 6.7 g, Glucose 20 g, Drop-out mix (3) 2.0 g,

Distilled water 1000 ml

(d) Gal-Ura-Leu medium

Bacto-yeast nitrogen base without amino acids 6.7 g,

Galactose 20 g Drop-out mix (3) 2.0 g, Distilled water 1000 ml Drop-out mix (1): Adenine 0.5 g, Lysine 2.0 g, Alanine 2.0 g, Methionine 2.0 g, Arginine 2.0 g, para-Aminobenzoic acid 0.2g, Asparagine 2.0 g. Phenylalanine 2.0 g, Aspartic acid 2.0 g, Proline 2.0 g, Cysteine 2.0 g, Serine 2.0 g, Glutamine 5 2.0 g, Threonine 2.0 g, Glutamic acid 2.0 g, Tryptophan 2.0 g, Glycine 2.0 g, Tyrosine 2.0 g, Histidine 2.0 g, Valine 2. 0 g, Inositol 2. 0 g, Isoleucine 2.0 g, Uracil 2.0 g, Leucine 10.0 g, Distilled water 1000 ml Drop-out mix (2): Dropout mix (1) except for leucine (10.0 g) Drop-out mix (3): Drop-out mix (1) except for uracil (2.0 g) and leucine (10.0 g) 10 (e) Glu-agar medium Solid medium in which 2%(W/V) agar is added to a medium (a) (f) Glu-Leu agar medium Solid medium in which 2% (W/V) agar is added to a medium (b) (g) Glu-Ura-Leu agar medium 15 Solid medium in which 2% (W/V) agar is added to a medium (c) (h) Gal-Ura-Leu agar medium Solid medium in which 2% (W/V) agar is added to a medium (d) Free text in Sequence Listing 20 SEQ ID NO:3 [0183] Designed oligonucleotide primer for PCR 25 SEQ ID NO:4 [0184] Designed oligonucleotide primer for PCR SEQ ID NO:5 30 [0185] Designed oligonucleotide primer for sequencing SEQ ID NO:6 35 [0186] Designed oligonucleotide primer for sequencing SEQ ID NO:7 [0187] Designed oligonucleotide primer for sequencing 40 SEQ ID NO:8 [0188] Designed oligonucleotide primer for sequencing 45 SEQ ID NO:9 [0189] Designed oligonucleotide primer for sequencing SEQ ID NO:10 50 [0190] Designed oligonucleotide primer for sequencing

[0191] Designed oligonucleotide primer for sequencing

SEQ ID NO:11

SEQ	10	NIC	 1
SEU	11.)	NI.	_

[0192] Designed oligonucleotide primer for sequencing

5 SEQ ID NO:15

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[0193] Designed oligonucleotide primer for PCR

SEQ ID NO:18

[0194] Designed oligonucleotide primer for PCR

SEQ ID NO:19

15 [0195] Designed oligonucleotide primer for PCR

SEQ ID NO:20

[0196] Designed oligonucleotide primer for sequencing

SEQ ID NO:21

[0197] Designed oligonucleotide primer for sequencing

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[0198] Designed oligonucleotide primer for sequencing

SEQ ID NO:23

[0199] Designed oligonucleotide primer for sequencing

SEQ ID NO:24

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SEQ ID NO:25

[0201] Designed oligonucleotide primer for sequencing

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[0202] Designed oligonucleotide primer for sequencing

45 SEQ ID NO:27

[0203] Designed oligonucleotide primer for sequencing

SEQ ID NO:28

[0204] Designed oligonucleotide primer for sequencing

SEQ ID NO:29

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[0218] Designed oligonucleotide primer for PCR

[0219] Designed oligonucleotide primer for DNA sequencing

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50	Ala Ile Thr Val Asn Asn Gln Thr Leu Pro Asp Thr Pro Asn Glu Leu
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55	Gly Ala Pro Ser Ala Phe Ala Asp Val Leu Thr Gly Ala Pro Ser Arg

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55	Leu	Thr	Lys	Lys	Ile	Glu	Val	Glu	Val	Gln	Gly	G1u	Ile	Ala	Ser	Leu

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				500	•			•	505					510		
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25		٠		580					585					590		·
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		690					695			-		700				
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	1105				1	110				1	115				1	120
55	Arg I	Leu /	Ala	Val	Lys	Ile	Leu	Glu	Lys	Tyr	His	His	Val	Val	Thr	Val

		1106		1100	- 405
5		1125		1130	1135
	Val Gly Asn	Gly Gln Glu	Ala Leu Asp	Ala Ile Lys	Glu Lys Arg Tyn
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10	Asp Val Ile I	Leu Met Asp	Val Gln Met	Pro Ile Met	Gly Gly Phe Glu
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	Thr Pro Ile	Ile Ala Leu	Thr Ala His	Ala Met Leu (	Gly Asp Arg Glu
20	1185	1190		1195	1200
	Lys Cys Ile (	Gln Ala Gln	Met Asp Glu	Tyr Leu Ser I	Lys Pro Leu Lys
		1205	1	210	1215
25	Gln Asn His I	eu Ile Gln	Thr Ile Leu	Lys Cys Ala 1	Thr Leu Gly Gly
	12	220	1225		1230
30	Ala Leu Leu (	Glu Lys Gly	Arg Glu Val	Arg Gln Ser A	Ala Asn Glu Glu
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	Thr Thr Thr G		Asn His Gly	•	Ser Pro Ser Leu
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	1 .		5					10		-			15		
30										•					
	gca tta			•									•		96
	Ala Leu	_	_	Asp	Leu	Pro		Thr	Asn	Val	Tyr		Asn	Lys	
35 -		2	0				25			•	٠	30			
		**			· :		•						-		4.4
40	ggg att						•							_	144
	Gly Ile	35	u Pro	GIY	Ala		Inr	Ala	GIU	Lys		Ala	Leu	GIU	
45	•	30				40					45				
70	cga gaa	rtt an	a acc	++0	σta	tee	200	at c		200		<b>722</b>	<b>700</b>		192
	Arg Glu I												•		192
50	50			204	55	-01	1 TO E		7111	60	Ten	274	117.0	*# B	
									•	30					

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5	Ala	Ile	Thr	Val	Asn	Asn	Ġ1n	Thr	Leu	Pro	Asp	Thr	Pro	Asn	Glu	<u>L</u> eu	•
	66	;				70					75					80	
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-				٠											•		
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	,			180				,	185	014	111 8	010	Leu	190	•	IILS	
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	Ile	Val	Thr	Ala	Val	Ala	Arg	Gly	Asp	Leu	Ser	Lys	Lys	Val	Gln	Ile	
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	225					230					235					240	
40			•						•								
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			•		Asp												
45					245		•			250				٠	255		
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	Val	Ala	Arg	Glu.	Val	Gly	Thr	<b>Gl</b> u	Gly	Ile	Leu	Gly	Gly	Gln	Ala	Lys	
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	Ile	Ser	Gly	Val	Asp	Gly	Thr	Trp	Lys	Glu	Leu	Thr	Asp	Asn	Va1	Asn	•
10			275					280					285				
													•				
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- 35	Gln	Phe	Ala	Arg	Glu	Va1	Thr	Lys	Ile	Ala	Arg	Glu	Va1	Gly	Thr	Glu	
-			435					440			·		445			•	
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	•															•	
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	Ile	Ser	Thr	Val	Thr	Gln	Ala	Ile	Ala	Asn	Gly	Asp	Met	Ser	Gla	Lys	
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	•																
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			675			-10 P		680		- ^^-	-, -		685			214	
		•															

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15	Tyr	Asn	Leu	Arg	Asp	Ser	I1e	Gln	Arg	Asn	Thr	Leu	Ala	Arg	G1u	Ala	
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25			•		725		•			730		•	•		735		
	•																
	cac	gag	att	cga	aca	cct	atg	aac	ggt	atc	att	ggt	atg	act	cag	ttg	225 <b>6</b>
30				Arg													
·		•		740	·				745					750			
35																	
-	aca	ctc	gac	acc	gat	ctt	act	caa	tat	caa	cga	gaa	atg	ctc	aac	att.	2304
				Thr													
40			755		-			760	•				765				
45	gīt	Cac	aac	ttg	gcc	aac	aøt.	tta	ttσ	acc	atc	att	oat	gat	att	ctc	2352
				Leu			•								•		
		770	******	Dea			775	Leu	Leu	4114	116	780	vab	лэр		Dea	
50		,,,					110	-				100					٠
	ga+	++0	+00			<b>7</b> 00			· a+	~*~	o+-	a+ a		~~~	-++		2400
	gat	LLA	LCA	aag	atc	gaa	gca	aac	cgt	atg	atc.	alg	gag	Rag	all <sub>.</sub>	CCA	2400

	Asp	Leu	Ser	Lys	Ile	Glu	Ala	Asn	Arg	Met	Ile	Met	Glu	Glu	Ile	Pro	
5	785					790					795		•			800	
					•											•	
10	tac	act	ctt	aga	gga	acc	gtc	ttc	aac	gco	ctc	aag	act	ctc	gct	gtc	2448
10	Tyr	Thr	Leu	Arg	Gly	Thr	Val	Phe	Asn	Ala	Leu	Lys	Thr	Leu	Ala	Val	
					805					810					815		
15																	
	aag	gca	aat	gag	aag	ttç	cta	gac	ctc	act	tac	cgc	gta	gat	agc	tca	2496
	Lys	Ala	Asn	Glu	Lys	Phe	Leu	Asp	Leu	Thr	Tyr	Arg	Val	Asp	Ser	Ser	
20				820					825					830			
				-				•				•					
25	gtt	cca	gat	cac	gtg	gtt	ggt	gat	.tca	ttc	cgt	ctt	cga	caa	gtt	att	2544
	Val	Pro	Asp	His	Val	Val	Gly	Asp	Ser	Phe	Arg	Leu	Arg	G1n	Val	Ile	
			835		• •			840					845				
30																	
•	ctc	aac	ttg	gtt	gga	aac	gct	atc	aag	ttc	aca	gag	cat	ggt	gaa	gtt	2592
35	Leu	Asn	Leu	Val	Gly	Asn	Ala	Ile	Lys	Phe	Thr	Glu	His	Gly	Glu	Va1	
-		850					855					860					
													-				
40	tcg	ttg	acc	atc	caa	aaa	gcc	gag	caa	gat	cat	tet	ECE	CCE	aac	<b>₽</b> aa	2640
				•	Gln									•			
45	865					870					875	-, -		•,•		880	
											0.0						, <del>-</del>
	***					<b></b>	_4.4			4						4.4	0000
50					ttt								•				2688
	Tyr	Ala	val	Glu		Cys	Val	Ser			GLy	lie	Gly	Ile		Ala	•
					885					890					895		
55																	•

5	gat	aag	cto	aat	ttg	att	tto	gac	act	ttc	cas	caa	gct	gac	gga	tct	2736
	Asp	Lys	Leu	ı Asn	Leu	Ile	Phe	Asp	Thr	Phe	Glm	Gln	Ala	Asp	G1y	Ser	
				900	•				905	;				910			
10																	
	atg	acg	agg	aaa	ttc	gga	ggt	act	ggt	cta	ggt	cta	tca	att	tcg	aag	2784
15	Met	Thr	Arg	Lys	Phe	G1y	Gly	Thr	Gly	Leu	Gly	Leu	Ser	Ile	Ser	Lys	
			915	i				920					925				
				•													
20	aga	ctt	gta	aac	ctc	atg	cgt	gga	gat	gtt	tgg	gtt	aag	agt	cag	tac	2832
				Asn													
25		930					935	-	-		•	940	_•-			-,-	
												• • •					
	gga	aaa	ggc.	agt	tca	ttc	tac.	ttc	200	t at	200	ait c	640	o <b>t</b> o	<b>700</b>	222	2000
30				Ser													2880
	945	275	01)	561	Per	950	1 9 2	LHe	1111	Cys		191	WLE	reu	AIS		÷
35	J-10	•				300					955					960 -	
	***		-4-														
				agt										•			2928
40	Ser	Asp	He	Ser		Ile	Gln	Lys	Gln		Lys	Pro	Tyr	Gln	Gly	His	
				•	965	•	•			<b>97</b> 0					975		
						٠.									٠	•	
45	aat	gtt	ttg	ttt	atc	gac	aaa	gga	cag	act	ggc	cat	ggc	aaa	gaa	ata	2976
	Asn	Val	Leu	Phe	Ile	Asp	Lys	Gly	Gln	Thr	Gly	His	Gly	Lys	Glu	Ile	
50				980					985					990,			
	atc	act	atg	ctt	aca	caa	ctt	ggt	ttg	gta	ccc	gtt	gtt	gtt	gac	tct	3024
i5																	

	Ile Th	r met	Leu	Thr	Gln	Leu	Gly	Leu	Val	Pro	Val	Val	Val	Asp	Ser	
5		995				•	1000					1005				
												٠			٠.	
10	gag ca	g cac	act	att	ctt	ctc	ggc	aat	gga	aga	acc	aag	gag	aag	att	3072
	Glu Gl	n His	Thr	Ile	Leu	Leu	Gly	Asn	Gly	Arg	Thr	Lys	Glu	Lys	Ile	
	101	0		•		1015			•		1020					
15																
	gct to	a acț	tat	gac	gtg	att	gtt	gtg	gac	tca	att	gag	tcc	gct	cga	3120
20	Ala Se	r Thr	Tyr	Asp	Val	Ile	Val	Val	Asp	Ser	Ile	Glu	Ser	Ala	Arg	
	1025			1	1030				:	1035				:	1040	
25	• •															
25	aaa ct	g cga	tca	atc	gat	gag	ttc	aag	tat	att	cca	att	gtt	ctc	tta	3168
	Lys Le	u Arg	Ser	Ile	Asp	Glu	Phe	Lys	Tyr	Ile	Pro	Ile	Val	Leu	Leu	
30	•		- 1	1045				]	1050				:	1055		
-		٠													•	
. 35	gct cc							٠.								3216
35	gct ccc	o Val	Ile					٠.				Asp	Leu			3216
35		o Val					Leu	٠.				Asp				3216
35		o Val	Ile				Leu	Lys				Asp	Leu			3216
***	Ala Pro	Val	Ile 1060 atg	His	Val act	Ser	Leu	Lys 1065 tta	Ser	Ala	Leu	Asp	Leu 1070 ggc	Gly	Ile ggt	3216 3264
***	Ala Pro	Val	Ile 1060 atg	His	Val act	Ser cca Pro	Leu tgt Cys	Lys 1065 tta	Ser	Ala	Leu gat Asp	Asp ctt Leu	Leu 1070 ggc	Gly	Ile ggt	
40	Ala Pro	Val	Ile 1060 atg	His	Val act	Ser cca Pro	Leu	Lys 1065 tta Leu	Ser acg Thr	Ala	Leu gat Asp	Asp	Leu 1070 ggc	Gly	Ile ggt	
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40	act too	t tac Tyr 1075	Ile 1060 atg Met	His acc Thr	Val act Thr	cca Pro	tgt Cys 1080	Lys 1065 tta Leu gct	ser acg Thr	Ala atc Ile	gat Asp	ctt Leu 085	Leu 1070 ggc Gly	Gly aat Asn gac	ggt Gly	
40	act to	t tac Tyr 1075	Ile 1060 atg Met	His acc Thr	Val act Thr gag Glu	cca Pro	tgt Cys 1080	Lys 1065 tta Leu gct	ser acg Thr	Ala atc Ile ccc Pro	gat Asp	ctt Leu 085	Leu 1070 ggc Gly	Gly aat Asn gac	ggt Gly	3264

5	ace aaa tco	ttc gac att	ctc ttg gcc	gaa gat aac	atc gtc aat	caa 3360
	Thr Lys Ser	Phe Asp Ile	Leu Leu Ala	Glu Asp Asn	Ile Val Asn (	Gln
10	1105	1110		1115	11	120
	,					•
	cgc tta gcg	gtg aag att	cta gaa aag	tat cac cac	gtc gtc aca g	gtc 3408
15	Arg Leu Ala	Val Lys Ile	Leu Glu Lys	Tyr His His	Val Val Thr V	/al
		1125		1130	1135	•
20				·		
	gtt ggc aat	ggt caa gaa	gca cta gat	gct atc aag	gag aaa cga t	ac 3456
25	Val Gly Asn	Gly Gln Glu	Ala Leu Asp	Ala Ile Lys	Glu Lys Arg T	îyr
25		1140	1145		1150	
30					gga gga ttc g	
-		Leu Met Asp	Val Gln Met	Pro Ile Met	Gly Gly Phe G	lu
35	1155	, .	1160	1	.165	
	-					
	·				gga acg caa a	
40	1170				Gly Thr Gln A	rg .
	. 1170	J	1175	1180		
45	700 001 017				· ,	2600
		•			ggt gat cgc g	
	1185	1190	inr Ala His		Gly Asp Arg G	
50	. 1100	1130		1195	12	uu
	aaa tgt att	caa gee caa	ato gat gaa	tat ctt tct	aag cct ctg a	aa 3648
55		804 246	5 800			,

	Lys	Cys	Ile	Gln	Ala	Gln	Met	Asp	Glu	Tyr	Leu	Ser	Lys	Pro	Leu	Lys	•
5	•				1205					1210					1215		
							-										٠
	caa	aat	cat	ctt	att	cag	acg	atc	ttg	aaa	tgt	gca	acc	ctt	gga	ggt	3696
10	Gln	. Asn	His	Leu	Ile	Gln	Thr	Ile	Leu	Lys	Cys	Ala	Thr	Leu	Gly	Gly	
				1220					1225					1230			•
15		٠															
,	gça	ttg	ctc	gag	aag	ggt	agg	gag	gtt	agg	caa	tcc	gct	aat	gaa	gag	3744
20	Ala	Leu	Leu	Glu	Lys	Gly	Arg	Glu	Val	Arg	Gln	Ser	Ala	Asn	Glu	Glu	
20			1235				Ċ	1240				į	1245			,	
25	agc	ccc	aat	tcg	caa	aat	ggt	cct	cgc	ggt	aca	cag	cat	cct	gca	tca	3792
	Ser	Pro	Asn	Ser	Gln	Asn	Gly	Pro	Arg	Gly	Thr	G1n	His	Pro	Ala	Ser	
30	1	L250					1255			٠	•	1260					
30	•									•							
•	agt	ccc	aca	cça	gcc	cat	atg	aga	ccg	gct	atc	gaa	cct	cgt	gça	tac	3840
35	Ser	Pro	Thr	Pro	Ala	His	Met	Arg	Pro	Ala	Ile	Glu	Pro	Arg	Ala	Tyr	
	1265	5			٠:	1270				1	275				:	1280	
40																•	
	acg	acc	act	ggc	cct	ata	ast	cat	gga	agt	gca	gag	agt	cct	tca	ctt	3888
	Thr	Thr	Thr	Gly	Pro	Ile	Asn	His	Gly	Ser	Ala	Glu-	Ser.	Pro	Ser	Leu	
15			•	1	285				1	290				• 1	1295		
						٠									0.		
50	gta	acg	gca	gat	gct	gag	gat	cca	ctt	gcg	agg	ctt	cta	atg	cgt	gcg	3936
	Val	Thr	Ala	Asp	Ala	Glu	Asp	Pro	Leu	Ala	Arg	Leu	Leu	Met	Arg	Ala	
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5	cat agc agc tag	3948
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# oligonucleotide primer for PCR

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30		
	<400> 5	÷
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_		•
40		•
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	<211> 30	
45	<212> DNA	
	<213> Artificial Sequence	
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# oligonucleotide primer for sequencing

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	(400) 6	
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	·	
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	<211> 30	
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	<213> Artificial Sequence	
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30	oligonucleotide primer for sequencing	
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35	acagaaggta ttctcggtgg acaagccaag	30
40	⟨210⟩ 8	
	· •	
45	<211> 30 -	
	<212> DNA	
	<213> Artificial Sequence	
50	·	
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# oligonucleotide primer for sequencing

5		
	< <b>400&gt;</b> 8	٠
10	gctagggagg tcggtaccga aggtagactg	30
15	<210> 9	
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40		•
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# oligonucleotide primer for sequencing

5		
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10		
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-		
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# oligonucleotide primer for sequencing

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,,,						
	•			•		
15	<210> 13					
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	<213> Botry	otinia fucke	liana		:	
			• .			
25	<400> 13	•				
	Met Glu Asp	Ser Thr Ile	Ala His Thr	Thr Ala Ile I	Leu Gln Thr Leu	
30	1	5	•	10	15	
_	Ala Leu Ser	Ser Ile Asp	Leu Pro Leu	Thr Asn Val 1	Tyr Gly Asn Lys	
		20	25	•	30	
<b>35</b> -	Gly Ile Arg	Leu Pro Gly	Ala Asp Thr	Ala Glu Lys I	Leu Ala Leu Glu	
	35		40		45	
40	Arg Glu Leu	Ala Ala Leu	Val Ser Arg	Val Gln Arg I	eu Glu Ala Arg	
	50		55	60		
	Ala Ile Thr	Val Asn Asn	Gln Thr Leu	Pro Asp Thr F	Pro Asn Glu Leu	
45	.65	70		75	. 80	
	Gly Ala Pro	Ser Ala Phe	Ala Asp Val	Leu Thr Gly A	Ala Pro Ser Arg	
50		85		90	95	
	Ala Ser Lys	Ser Thr Thr	Ser Arg-Gln	Gln Leu Val A	isn Ser Leu Leu	•
		100	105	•	110	

73

	Ala	Ala	Arg	Glu	Ala	Pro	Thr	Gly	Gly	G1u	Arg	Pro	Pro	Lys	Phe	Thr
5			115					120				-	125			
	Lys	Leu	Ser	Asp	Glu	Glu	Leu	Glu	Ala	Leu	Arg	Glu	His	Val	Asp	His
	-	130					135					140				
10	Gln	Ser	Lys	Gln	Leu	Asp	Ser	Gln	Lys	Ser	Glu	Leu	Ala	Gly	Val	His
	145					150					165					160
15	Ala	G1n	Leu	Phe	Glu	Gln	Lys	Gln	Arg	Gln	Glu	Gln	Ala	Leu	Asn	Val
					165					170					175	
	Leu	Glu	Val	Glu	Arg	Va1	Ala	Ala	Leu	Glu	Arg	Glu	Leu	Lys	Lys	His
20 .				180					185					190		
	Gln	Gln	Ala	Asn	Glu	Ala	Phe	Gln	Lys	Ala	Leu	Arg	Glu	Ile	Gly	Glu
25			195					200			٠.		205			
	Ile	Va1	Thr	Ala	Va1	Ala	Arg	G1y	Asp	Leu	Ser	Lys	Lys	Val	Gln	Ile
20		210					215					220	•			
30	His	Ser	Val	G1u	Met	Asp	Pro	Glu	Ile	Thr	Thr	Phe	Lys	Arg	Val	Ile
•	225			Δ	•	230					235		-			240
35	Asn	Thr	Met	Met	Asp	Gln	Leu	Gln	Ile	Phe	Ser	Ser	G1u	.Va1	Ser	Arg
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40 .	Val	Ala	Arg	Glu	Val	Gly	Thr	Glu	G1y	Ile	Leu	Gly	Gly	<b>G</b> ln	Ala	Lys
			•	260					265					270		
	Ile	Ser		Va1	Asp	G1y	Thr	Trp	Lys	Glu	Leu	Thr	Asp	Asn	Val	Asn
45			275					280					285	·		
	Val		Ala	Gln	Asn	Leu	Thr	Asp	Gln.	Val	Arg	Glu	Ile	Ala	Ser	Val
50		290					295					300	•			
			Ala	Val	Ala			Asp	Leu	Thr	•	•	Ile	Glu	Arg	Pro
	305					310	)				315	i .				320

	Ala	Gln	Gly	Glu	Ile	Leu	Gln	Leu	Gln	Gln	Thr	Tle	Asn	Thr	Met	Val
5					325					330	•		•		335	;
	Asp	Gln	Leu	Arg	Thr	Phe	Ala	Ala	Glu	·Val	Thr	Arg	Val	Ala	Arg	Asp
				340					345					<b>35</b> 0		
10	Val	Gly	Thr	Glu	Gly	Ile	Leu	Gly	Gly	Gln	Ala	Glu	Ser	Glu	G1y	Val
•			355					360					365			
15	Gln	Gly	Met	Trp	Asn	Thr	Leu	Tle	Val	Asn	Val	Asn	Ala	Met	Ala	Asn
		370					375					380				
20	Asn	Leu	Thr	Thr	GIn	Val	Arg	Asp	Ile	Ala	Ile	Val	Thr	Thr	Ala	Val
	385					<b>39</b> 0			٠		395					400
	Ala	Lys	Gly	Asp	Leu	Thr	Gln	Lys	Val	Gln	Ala	Glu	Cys	Lys	Gly	Glu
25					405					410					415	
	Ilė	Lys	G1n	Leu	Lys	G1u	Thr	Ile	Asp	Ser	Met	Val	Asp	G1n	Leu	Gln
30				420					425					430		
	G1n	Phe	Ala	Arg	Glu	Val	Thr	Lys	Ile	Ala	Arg	Glu	Val	G1y	Thr	Glu
	•		435	•				440					445			
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		450					455					460				
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	465				•	470					475					480
	Gln	Val	Arg	G1u	Ile	Ala	Lys	Val	Thr	Thr	Ala	Val	Ala	Arg	G1y	Asp
45					485					490					495	
	Leu	Thr	Lys	Lys	Ile	Glu	Val	Glu	Val	G1n	Gly	G1u	Ile	Ala	Ser	Leu
50				500					505					510		
	Lys	Asp		Ile	Asn	Thr	Met		Asp	Arg	Leu	Ser	Thr	Phe	Ala	Phe
			515			•		520					525		•	

	Glu	Val	Ser	Lys	Val	Ala	Arg	G1 u	Val	Gly	Thr	Asp	Gly	Thr	Leu	Gly
5		530					535					540				
	G1y	G1n	A1a	Gln	Val	Asp	Asn	Val	G1u	Gly	Lys	Trp	Lys	Asp	Leu	Thr
10	545					550		•			555					560
,,,	Glu	Asn	Va1	Asn	Thr	Met	Ala	Arg	Asn	Leu	Thr	Thr	Gln	Val	Arg	Gly
	-			•	565					570					<b>57</b> 5	
15	Ile	Ser	. Thr	Val	Thr	Gln	Ala	Ile	Ala	Asn	Gly	Asp	Met	Ser	Gln	Lys
				580					585					590		
20	Ile	Glu	Val	Ala	Ala	Ala	Gly	Glu	Ile	Leu	Ile	Leu	Lys	Glu	Thr	Ile
			595					600					605			
	Asn	Asn	Met	Val	Asp	Arg	Leu	Ser	Ile	Phe	Ser	Asn	Glu	Va1	Gln	Arg
?5		610	•			٠	615				•	620			• .	
	Val	Ala	Lys	Asp	Val	Gly	Val	Asp	Gly	Lys	Met	Gly	Gly	G1n	Ala	Asp
10	625					630					635				•	640
	Val	Ala	Gly	Ile	Gly	Gly	Arg	Trp	Lys	Glu	Ile	Thr	Thr	Asp	Val	Asn
					645	•	•			650					655	
-	Thr	Met	Ala	Asn	Asn	Leu	Thr	Thr	Gln	Val	Arg	Ala	Phe	Gly	Asp	Ile
				660					665					670		
o	Thr	Asn <sub>.</sub>	Ala	Ala	Thr	Asp	Gly	Asp	Phe	Thr	Lys	Leu	Ile	Thr	Val	Glu
			675					680	,		• .		685			
	Ala	Ser	G1y	Glu	Met	Asp	Glu	Leu	Lys	Arg	Lys	Ile	Asn	Gln	Met	Val
<b>.</b>		690					695					700				
	Tyr	Asn	Leu	Arg	Asp	Ser	Ile	Gln	Arg	Asn	Thr	Leu	Ala	Arg	Glu	Ala
0	705	•				710					715					720
	Ala	Glu	Phe	Ala	Asn	Arg	Thr	Lys	Ser	Glu	Phe	Leu	Ala	Asn	Met	Ser
					725					730			•		735	

	His	Glu	Ile	Arg	Thr	Pro	Met	Asn	Gly	Ile	Ile	Gly	Met	Thr	Gln	Leu
5				740					745					750		
	Thr	Leu	Asp	Thr	Asp	Leu	Thr	Gln	Tyr	Gln	Arg	Glu	Met	Leu	Asn	Ile
			755	٠				760					765	,		•
10	Val	His	Asn	Leu	Ala	Asn	Ser	Leu	Leu	Thr	Ile	Ile	Asp	Asp	Ile	Leu
		770		•		•	775					780				
15	Asp	Leu	Ser	Lys	Ile	Glu	Ala	Asn	Arg	Met	Ile	Met	Glu	Glu	Ile	Pro
	785					790					795					800
	Tyr	Thr	Leu	Arg	Gly	Thr	Val	Phe	Asn	Ala	Leu	Lys	Thr	Leu	Ala	Val
20					805					810					815	
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?5				820					825					830		
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		850		•			855					860				
35	Ser	Leu	Thr	Ile	Gln	Lys	Ala	G1u	Gln	Asp	His	Cys	Ala	Pro	Asn	Glu
-	865		•			870					875			•		880
	Tyr	Ala	Val	Glu	Phe	Cys	Val	Ser	Asp	Thr	G1y	Ile	Gly	Ile	Gln	Ala
10					885					890	•				895	
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5				900					905					910		
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	•		915					920					925			
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		930					935					940			•	

	Gly	Lys	Gly	Ser	Ser	Phe	Tyr	Phe	Thr	Cys	Thr	Val	Arg	Leu	Ala	Thr
5	945					950					955					960
	Ser	Asp	Ile	5er	Phe	Ile	Gln	Lys	Gln	Leu	Lys	Pro	Tyŗ	Gln	Gly	His
10					965					970					975	
·	Asn	Val	Leu	Phe	Ile	Asp	Lys	Gly	Gln	Thr	Gly	His	Gly	Lys	Glu	Ile
15				980	).				985					990		
15	Ile	Thr	Met	Leu	Thr	G1n	Leù	G1y	Leu	Val	Pro	Val	Val	Val	Asp	Ser
			995					1000					1005			
20	Glu (	Gln	His	Thr	Ile	Leu	Leu	Gly	Asn	Gly	Arg	Thr	Lys	Glu	Lys	Ile
-	1	010				1	1015			•	]	1020		••		
25	Ala	Ser	Thr	Tyr	Asp	Val	Ile	Val	Val	Asp	Ser	Ile	G1u	Ser.	Ala	Arg
	1025				1	1030				1	1035				. 1	1040
	Lys !	Leu	Arg	Ser	Ile	Asp	Glu	Phe	Lys	Tyr	Ile	Pro	Ile	Val	Leu	Leu
30				1	1045				1	050				1	1055	
	Ala I	Pro	Val	Ile	His	٧al	Ser	Leu	Lys	Ser	Ala	Leu	Asp	Leu	Gly	Ile
35			1	060				1	065				1	1070		
-	Thr S	Ser	Tyr	Met	The	The second	_	<u> </u>	-	TL-		Acn			A	C1
			-/-		1111	ınr	Pro	Cys	Leu	inr	lle	vəh	Leu	Gly	Asn	.GLY
			.075		1111	ınr		.080	Leu	inr.	ile		Leu .085	Gly	ASN	.GTA
40	Met ]	1	075				1	.080				1	.085			
40	Met ]	1	075			Glu	1	.080			Pro	1	.085			
40	Met ]	1 []e 090	075 Pro	Ala	Leu	Glu I	1 Asn 095	.080 Arg	Ala	Ala	Pro 1	1 Ser 100	.085 Leu	Ala	Asp	Asn
	Met ]	1 []e 090	075 Pro	Ala	Leu Asp	Glu I	1 Asn 095	.080 Arg	Ala	Ala Glu	Pro 1	1 Ser 100	.085 Leu	Ala	Asp Asn	Asn
45	Met 1	1   1e   090   Lys	075 Pro Ser	Ala Phe	Leu Asp	Glu I Ile 110	Asn 095 Leu	Arg Leu	Ala Ala	Ala Glu 1	Pro 1 Asp 115	Ser 100 Asn	.085 Leu Ile	Ala Val	Asp Asn 1	Asn Gln 120
	Met 1 1( Thr 1 1105	1   1e   090   Lys	075 Pro Ser	Ala Phe Val	Leu Asp	Glu I Ile 110	Asn 095 Leu	Arg Leu	Ala Ala Lys	Ala Glu 1	Pro 1 Asp 115	Ser 100 Asn	.085 Leu Ile	Ala Val	Asp Asn 1	Asn Gln 120
45	Met 1 1( Thr 1 1105	1 Ile 090 Lys Leu	O75 Pro Ser Ala	Ala Phe Val	Leu Asp 1 Lys	Glu I Ile 110 Ile	Asn 095 Leu Leu	.080 Arg Leu Glu	Ala Ala Lys	Ala Glu 1 Tyr 130	Pro 1 Asp 115 His	Ser 100 Asn His	.085 Leu Ile Val	Ala Val Val	Asp Asn 1 Thr	Asn Gln 120 Val

11	.55		1160		1165	
Ala Thr A	la Lys Ile	Arg Glu	Tyr Glu	Arg Ser	Leu Gly	Thr Gln Arg
1170		1175			1180	•
Thr Pro I	le Ile Ala	Leu Thr	Ala His	Ala Met	Leu Gly	Asp Arg Glu
1185		1190		1195		1200
Lys Cys I	le Gln Ala	Gln Met	Asp Glu	Tyr Leu	Ser Lys	Pro Leu Lys
	1205			1210		1215
Gln Asn H	is Leu Ile	Gln Thr	Ile Leu	Lys Cys	Ala Thr	Leu Gly Gly
	1220		1225		1.	230
Ala Leu L	eu Glu Lys	Gly Arg	Glu Val	Arg Gln	Ser Ala	Asn Glu Glu
12	35		1240		1245	•
Ser Pro A	sn Ser Gln	Asn Gly	Pro Arg	Gly Thr	Gln His	Pro Ala Ser
1250		1255		. 1	1260	
Ser Pro T	hr Pro Ala	His Met	Arg Pro	Ala Ile	Glu Pro	Arg Ala Tyr
1265	1	1270		1275		1280
The The T	hr Gly Pro	Ile Asn	His Gly	Ser Ala	Glu Ser	Pro Ser Leu
	1285					1296
Val Thr A		Glu Asp		Ala Arg	Leu Leu l	Wet Arg Ala
			1305		13	310
	•					
13)	15					
		•			٠	
(210) 14						
	Ala Thr A  1170 Thr Pro I  1185 Lys Cys I  Gln Asn H  Ala Leu L  12  Ser Pro A  1250 Ser Pro T  1265 Thr Thr T  Val Thr A  His Ser	Thr Pro Ile Ile Ala 1185 Lys Cys Ile Gln Ala 1205 Gln Asn His Leu Ile 1220 Ala Leu Leu Glu Lys 1235 Ser Pro Asn Ser Gln 1250 Ser Pro Thr Pro Ala 1265 Thr Thr Thr Gly Pro 1285 Val Thr Ala Asp Ala 1300 His Ser Ser	Ala Thr Ala Lys Ile Arg Glu  1170  1175  Thr Pro Ile Ile Ala Leu Thr  1185  1190  Lys Cys Ile Gln Ala Gln Met  1205  Gln Asn His Leu Ile Gln Thr  1220  Ala Leu Leu Glu Lys Gly Arg  1235  Ser Pro Asn Ser Gln Asn Gly  1250  Ser Pro Thr Pro Ala His Met  1265  1270  Thr Thr Thr Gly Pro Ile Asn  1285  Val Thr Ala Asp Ala Glu Asp  1300  His Ser Ser  1315	Ala Thr Ala Lys Ile Arg Glu Tyr Glu  1170  1175  Thr Pro Ile Ile Ala Leu Thr Ala His  1185  1190  Lys Cys Ile Gln Ala Gln Met Asp Glu  1205  Gln Asn His Leu Ile Gln Thr Ile Leu  1220  1225  Ala Leu Leu Glu Lys Gly Arg Glu Val  1235  1240  Ser Pro Asn Ser Gln Asn Gly Pro Arg  1250  1255  Ser Pro Thr Pro Ala His Met Arg Pro  1265  1270  Thr Thr Thr Gly Pro Ile Asn His Gly  1285  Val Thr Ala Asp Ala Glu Asp Pro Leu  1300  1305  Kis Ser Ser  1315	Ala Thr Ala Lys Ile Arg Glu Tyr Glu Arg Ser  1170 1175  Thr Pro Ile Ile Ala Leu Thr Ala His Ala Met 1185 1190 1195  Lys Cys Ile Gln Ala Gln Met Asp Glu Tyr Leu 1205 1210  Gln Asn His Leu Ile Gln Thr Ile Leu Lys Cys 1220 1225  Ala Leu Leu Glu Lys Gly Arg Glu Val Arg Gln 1235 1240  Ser Pro Asn Ser Gln Asn Gly Pro Arg Gly Thr 1250 1255  Ser Pro Thr Pro Ala His Met Arg Pro Ala Ile 1265 1270 1275  Thr Thr Thr Gly Pro Ile Asn His Gly Ser Ala 1285 1290  Val Thr Ala Asp Ala Glu Asp Pro Leu Ala Arg 1300 1305  His Ser Ser	Ala Thr Ala Lys Ile Arg Glu Tyr Glu Arg Ser Leu Gly 1170 1175 1180  Thr Pro Ile Ile Ala Leu Thr Ala His Ala Met Leu Gly 1185 1190 1195  Lys Cys Ile Gln Ala Gln Met Asp Glu Tyr Leu Ser Lys 1205 1210  Gln Asn His Leu Ile Gln Thr Ile Leu Lys Cys Ala Thr 1220 1225  Ala Leu Leu Glu Lys Gly Arg Glu Val Arg Gln Ser Ala 1235 1240 1245  Ser Pro Asn Ser Gln Asn Gly Pro Arg Gly Thr Gln His 1250 1255 1260  Ser Pro Thr Pro Ala His Met Arg Pro Ala Ile Glu Pro 1265 1270 1275  Thr Thr Thr Gly Pro Ile Asn His Gly Ser Ala Glu Ser 1285 1290  Val Thr Ala Asp Ala Glu Asp Pro Leu Ala Arg Leu Leu Il 1300 1305  His Ser Ser 1315

<211> 3948

	(2)	12> 1	DNA															
5	<21	13> I	Botry	rotin	ia f	ucke	lian	a								. <del>"</del>		
10	<22														•			
		21> C 22> (	DS (1)	(394	.R)													
15				(002	,													
	<40	0> 1	4					•										
20													ctg				4	48
	1		v vah	267	5		VIS	DIS	1111	10	via	116	Leu	GIN	15			
25										•								
													tac				. 9	<del>9</del> 6
30	AIS	Leu	5er	5er 20	ITe	Asp	Leu	Pro	Leu 25	Thr	Asn	Val	Тут	Gly 30	Asn	Lys		
	•					•							-					
35													ctt				14	14
	Gly	Ile	Arg 35	Leu	Pro	Gly	Ala	Asp 40	Thr	Ala	Glu	Lys	Leu 45	Ala	Leu	G1u		
40						•		30		•						÷		
	cga	gaa	ctt	g¢g	gcc	ttg	gta	tcc	aga	gtc	caa	aga	tta	gaa	gça	agg	19	2
45	Arg		Leu	Ala	Ala	Leu		Ser	Arg	Val	Gl'n		Leu	Glu	Ala	Arg		
		50					55			•		60						
50	gcg	atc	aca	gtc	aat	aat	caa	acc	ctg	ccc	gat	acg	ccg	aat	gaa	tta	24	0
E E	Ala	Ile	Thr	Val	Asn	Asn	Gln	Thr	Leu	Pro	Asp	Thr	Pro	Asn	Glu	Leu		

	65					70					75					80	
5																	
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10	Gly	Ala	Pro	Ser	Ala	Phe	Ala	Asp	Val	Leu	Thr	Gly	Ala	Pro	Ser	Arg	
		٠		•	. 85					90					95	_	٠
15	gcc	tca	aag	agt	act	aca	tcc	cga	caa	cag	ctc	gta	aat	tcg	ttg	ctt	336
	Ala	Ser	Lys	Ser	Thr	Thr	Ser	Arg	Glņ	Gln	Leu	Val	Asn	Ser	Leu	Leu	
20				100					105					110	•		
	gcc	gcc	aga	gaa	gcg	ccc	acc	ggc	ggt	gaa	aga	cct	cct	888	ttt	acg	384
<b>25</b> .	Ala	Ala	Arg	Glu	Ala	Pro	Thr	Gly	Gly	Glu	Arg	Pro	Pro	Lys	Phe	Thr	-
	•		115					120		•			125		•		
30 .	.•	•															
	aaa	tta	agt	gac	gag	gaa	ctc	gaa	gca	ctc	cgc	gaa	cat	gtc	gac	cat	432
	Lys	Leu	Ser	Asp	Glu	Glu	Leu	Glu	Ala	Leu	Arg	Glu	His	Va1	Asp	His	٠
35 -		130					135					140			•		
40	caa	tcg	aaa	caa	ctc	gat	agt	caa	aaa	tct	gag	ctg	gcc	ggt	gta	cat	480
	Gln	Ser	Lys	G1n	Leu	Asp	Ser	G1n	Lys	Ser	Glu	Leu	Ala	G1y	Val	His	
	145					150					155					160	•
<b>1</b> 5																	
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50					Glu											•	
			-		165		-		J	170					175		

5	· cti	t ga	a gto	c gas	a cgo	gta	a gc	a gct	tct	c ga	a ag	a gaa	a ct	g aa	g aa	g cat	576
	Lei	ı Glı	u Val	l Gli	ı Arg	y Val	Ala	a Ala	a Lei	u Gli	u Arı	g Glu	ı Lei	ı Ly:	s Ly:	s His	
				180	)				185	5				190	9		
10												•					
	caa	caa	a gcc	aac	gag	gct	tto	caa	aaa	gc1	t cta	a cgg	gaz	ata	a gga	gag	624
15	G1n	G1r	n Ala	Asn	Glu	Ala	Phe	Gln	Lys	Ala	a Leu	ı Arg	G1v	Ile	• Gl3	Glu	
			195	;				200	)		•	•	205	;		•	
		-						•									
20	att	gto	aca	gct	gta	gct	agg	ggt	gat	cto	agt	aag	aag	gta	caa	atc	672
	Ile	Va1	Thr	Ala	Val	Ala	Arg	Gly	Asp	Lev	Ser	Lys	Lys	Val	Gln	Ile	
25		210					215					220	ı				
	•							•					. <i>.</i>				•
	cac	tcc	gtg	gag	atg	gac	cct	gag	att	aca	act	ttc	aag	cgt	gtt	att	720
30	His	Ser	Val	Glu	Met	Asp	Pro	Glu	Ile	Thr	Thr	Phe	Lys	Arg	Val	Ile	
	225					230		-			235					240	
35						-											
	aat	act	atg	atg	gat	caa	ctt	cag	ata	ttc	tct	agt	gag	gtt	tct	cgt	768
	Asn	Thr	Met	Met	Asp	G1n	Leu	Gln	Ile	Phe	Ser	Ser	Glu	Va1	Ser	Arg	•
<b>4</b> 0					245					250		•		٠.	255		
							÷							•			•
<b>4</b> 5	gta	gct	aga	gag	gtc	ggc	aca	gaa	ggt	att	ctc	ggt	gga	caa	gcc	aag	816
	Val	Ala	Arg	Glu	Val	Gly	Thr	Glu	Gly	Ile	Leu	Gly	Gly	Gln	Ala	Lys	
				260					265				-	270			
50									٠								•
	att	tct	ggt	gtt	gat	ggt	aca	tgg	aag	gag	ttg	act	gac	aat	gtc	aac	864
55	Ile	Ser	G1y	Val	Asp	Gly	Thr	Trp	Lys	Glu	Leu	Thr	Asp	Asn	Val	Λsn	

5			275					280					285				
10		atg	Ala				Thr					Glu		_		-	912
15	act	290		gt a	***	ont.	295	~~*	***			300	_**				
20	Thr	act Thr				His					Gln					Pro	960
20	305		, 			310		•			315	_				320	
25		Gln			Ile					Gln					Met		1008
30	•				325					330					335		
 35	•	caa Gln							•								1056
	·			340					345					350			
40		gga Gly	٠.		•												1104
45			355					360					<b>36</b> 5				·
50		ggc Gly															1152
55		370					375					380					

5	aac	ctc	acc	aca	caa	gtg	cgc	gat	ata	gcc	att	gtc	aca	aca	gct	gtc	1200
•	Asn	Leu	Thr	Thr	Gln	Val	Arg	Λsp	Ile	Ala	Ile	Val	Thr	Thr	Ala	Val	
	385		-			390		٠			395					400	
10																	
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15	Ala	Lys	Gly	Asp	Leu	Thr	Gln	Lys	Val	Gln	Ala	Glu	Cys	Lys	Gly	Glu	
					405					410					415		
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	Ile	Lys	Gln	Leu	Lys	Glu	Thr	Ile	Asn	Ser	Met	Yal	Asp	Gln	Leu	G1n	
25			-	420					425	·		•		430			•
	caa	ttt	gcg	cga	gaa	gtc	acg	aag	att	gct	agg	gag	gtc	ggt	acc	gaa	1344
30	Gln	Phe	Ala	Arg	Glu	Val	Thr	Lys	Ile	Ala	Arg	Glu	Val	Gly	Thr	Glu	
			435				•	440		-			445		•		
35				·													
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	Gly	Arg	Leu	Gly	Gly	Gln	Ala	Thr	Val	His	Asp	Va1	Glu	Gly	Thr	Trp	
10		450					455					460					
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	Arg	Asp	Leu	Thr	G1u	Asn	Val	Asn	Gly	Met	Ala	Met	Asn	Leu	Thr	Thr	
	465			-	•	470					475					480	
io																	
	caa	gta	cga	gag	att	gca	aag	gtt	acc	acc	gct	gtc	gcc	aga	gga	gat	1488
5				Glu							•						
-																	

5					48	5				490	)				498	5	
	tt	g a	c a	ag aa	g atı	c gaa	gto	gag	gtt	; çag	gg8	gae	atc	gct	tcg	ctg	1536
10	Le	u Tì	ur L	ys Ly	s Ile	• G1u	ı Val	Glu	Val	Gln	G1y	Glu	ı Ile	Ala	Ser	Leu	
				50	0				505	;				510	ı		
15	aa	a. <b>g</b> z	t a	c at	c aac	e acc	ato	ata	aec	9073		2 mt	200	***	<b></b>	***	1504
					e Asn												1584
20			51				mo t	520		n. 5	, Leu	361	525	rne	VIS	Life	
25	ga	ggt	t ag	c aa	a gtc	gcc	agg	gag	gtc	gga	act	gat	ggg	act	ctt	ggt	1632
	Gla	ı Va	l Se	r Ly	s Val	Ala	Arg	Glu	Val	Gly	Thr	Asp	Gly	Thr	Leu	Gly	
		53	0				535					540					
30																	
	gga	ca	a go	g ca	gtt	gat	aac	gtc	gaa	gga	aag	tgg	aaa	gac	ctc	act	1680
35	Gly	G1	n Al	a Gli	ı Val	Asp	Asn	Val	Glu	Gly	Lys	Trp	Lys	Asp	Leu	Thr	
	545	· .				550					5 <b>55</b>	•			•	560	•
40					acc			•									1728
	Glu	Ası	n Va	l Asr	Thr	Met	Ala	Arg	Asn	Leu	Thr	Thr	Gln	Val	Arg	Gly	•
45		•	•	•	565					570				٠	575		
				0.									ė				•
50					aca												1776
	lie	261	Th		Thr	Gln	Ala			Asn	Gly	Asp			Gln	Lys	
	·			580					<b>58</b> 5					590			
55																	•

5	ati	t gag	gtt	gct	gct	gcg	ggt	gaa	ata	ctc	ata	cta	aag	gaa	acc	ata	1824
	Ile	Glu	Val	Ala	Ala	Ala	Gly	Glu	Ile	Leu	Ile	Leu	Lys	Glu	Thr	Ile	
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10														•			
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15	Asn	ı Asn	Met	Val	Asp	Arg	Leu	Ser	Ile	Phe	Ser	Asn	Glu	Val	Gln	Arg	•
	•	610	)				615					620					·.
						•		•		٠							
20	gto	gcc	aaa	gat	gtg	ggt	gtg	gat	ggt	aag	atg	ggt	ggc	caa	gct	gac	1920
	Val	Ala	Lys	Asp	Val	G1ÿ	Val	Asp	Gly	Lys	Met	Gly	Gly	Gln	Ala	Asp	
<b>25</b>	625	;				630					635		•			640	
		•															
	gtt	gct	ggg	att	ggc	ggc	cgt	tgg	aaa	gag	atc	aca	acg	gat	gtc	aat	1968
30	Val	Ala	Gly	Ile	Gly	Gly	Arg	Trp	Lys	Glu	Ile	Thr	Thr	Asp	Va1	Asn	
					645					650				•	655		
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-	acc	atg	gct	aac	aac	ttg	aca	acc	caa	gtg	cgc	gcc	ttt	ggt	gat	ata	2016
	Thr	Met	Ala	Asn	Asn	Leu	Thr	Thr	Gln	Val	Arg	Ala	Phe	G1y	Asp	Ile	
40				660					665					670	-		
															•		
45	act	aac	gcc	gça	acc	gat	ggc	gac	ttc	aca	aaa	ttg	atc	act	gtc	gag	2064
	Thr	Asn	Ala	Ala	Thr	Asp	Gly	Asp	Phe	Thr	Lys	Leu	Ile	Thr	Val	Glu	
			675					680					685				
50			•					-									
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		•															
5		690					695					700	•				
			•														
10	tac	aat	ctg	agg	gac	agt	att	caa	aga	aac	acc	ttg	gct	agg	gag	gct	2160
	Tyr	Asn	Leu	Arg	Asp	Ser	Ile	Gln	Arg	Asn	Thr	Leu	Ala	Arg	Glu	Ala	
	705	;				710					715					720	
15				٠													
•	gcc	gaa	ttc	gcc	aat	agg	acg	aag	tct	gaa	ttc	ttg	gct	aac	atg	tct	2208
20	Ala	Glu	Phe	Ala	Asn	Arg	Thr	Lys	Ser	Glu	Phe	Leu	Ala	Asn	Met	Ser	
				•	725					730			•		735		
			-														•
25	cac	gag	att	cga	aca	cct	atg	aac	ggt	atc	att	ggt	atg	act	cag	ttg	2256
	His	Glu	Ile	Arg	Thr	Pro	Met	Asn	Gly	Ile	Ile	Gly	Met	Thr	Gln	Leu	
30		-	•	740					745			·		750			
									•								
	aca	ctc	gac	acc	gat	ctt	act	caa	tat	caa	cga	gaa	atg	ctc	<b>aa</b> c	att	2304
35 -	Thr	Leu	Asp	Thr	Asp	Leu	Thr	Gln	Tyr	Gln	Arg	G1u	Met	Leu	Asn	Ile	
			<b>75</b> 5		•			760				'	765				
40								٠									
	gtt	cac	aac	ttg	gcc	aac	agt	tta	ttg	acc	atc	att	gat	gat	att	ctc	2352
	Val	His	Asn	Leu	Ala	Asn	Ser	Leu	Leu	Thr	Ile	Ile	Asp	Asp	Ile	Leu	
45		770					775					780					
			•										•				. •
50	gat	tta	tca	aag	atc	gaa	gca	aac	cgt	atg	atc	atg	gag	gag	att	cca	2400
	Asp	Leu	Ser	Lys	Ile	G1u	Ala	Asn	Arg	Met	Ile	Met	Glu	Glu	Ile	Pro	
	785					790			•		795					800	•
<b>55</b> .																	

5	tac	act	ctt	aga	gga	acc	gto	ttc	aac	gco	ctc	aag	act	CTC	gut	gtc	2448
	Тул	Thr	Lei	Arg	Gly	Thr	Val	Phe	Asn	Ala	Leu	Lys	Thr	Leu	Ala	Val	
					805					810					815	;	
10																	
	aag	gca	aat	gag	aag	ttc	cta	gac	ctc	act	tac	cgc	gta	gat	agc	tca	2496
15	Lys	Ala	Asn	Glu	Lys	Phe	Leu	Asp	Leu	Thr	Tyr	Arg	Val	Asp	Ser	Ser	
				820					825					830			
																·	
20	gtt	cca	gat	cac	gtg	gtt	ggt	gat	tca	ttc	cgt	ctt	cga	caa	gtt	att	2544
	Val	Pro	Asp	His	Val	Val	Gly	Asp	Ser	Phe	Arg	Leu	Arg	Gln	Va1	Ile	
25			835					840					845				
	ctc	aac	ttg	gtt	gga	aac	gct	atc	aag	ttc	aca	gag	cat	ggt	gaa	gtt	2592
30	Leu	Asn	Leu	Val	Gly	Asn	Ala	Ile	Lys	Phe	Thr	Glu	His	Gly	Glu	Val	
		850					855					860					•
35									÷				•				
-	tcg	ttg	acc	atc	caa	aaa	gcc	gag	caa	gat	cat	tgt	gcg	ccg	aac	gaa	2640
	Ser	Leu	Thr	Ile	Gln	Lys	Ala	G1u	Gln	Asp	His	Cys	Ala	Pro	Asn	Glu	•
40	865	-				870					875				٠.	880	,
											•					•	
45	tat	gca	gtc	gag	ttt	tgt	gtt	tct	gac	act	ggt	atc	ggt	atc	caa	gct	2688
	Tyr	Ala	Va1	-G1u	Phe	Cys	Va1	Ser	Asp	Thr	Gly	Ile	Gly	Ile	Gln	Ala	
				•	885			÷		890					895		
50																	
	gat	aag	ctc	aat	ttg	att	ttc	gac	act	ttc	caa	caa	gct	gac	gga	tct	2736
55	Asp	Lys	Leu	Asn	Leu	Ile	Phe	Asp	Thr	Phe	Gln	G1n	Ala	Asp	Gly	Ser	

5				900					905					910			
10											ggt Gly					aag Lys	2784
15			915					920					925				
											tgg Trp						2832
20		930					935	•				940				,	
25							•				acc						2880
30	945	Lys	GIÀ	ser	261	950	ıyr	Pne	ınr	Cys	Thr 955	vai	Arg	Leu	Ala	960	
	tca	gat	atc	agt	<b>t</b> tc	att	cag	<b>aaa</b>	caa	ctc	aag	cca	tat	caa	ggt	cac	2928
35 -	Ser	Asp	Ile	Ser	Phe 965	Ile	Gln	Lys	Gln	Leu 970	Lys	Pro	Tyr	G1n	G1y 975	His	
40	aat	gtt	ttg	ttt	atc	gac	aaa	gga	cag	act	ggc	cat	ggc	8aa	gaa	ata	2976
											Gly				•		
45																	
50											ccc Pro			_	_		3024
			995				1	000				. 1	005			٠	

_	gag	g cag	cac	act	att	ctt	ctc	ggo	aat	ggs	aga	acc	aag	g gag	g aag	att	3072
5	Glu	ı Gln	ı His	Thr	· Ile	Leu	Leu	Gly	Asr	ı Gly	Arg	Thr	Lys	s Glı	ı Lys	Ile	
	_	1010	)				1015	i				1020	)				
10																	
	gct	tca	act	tat	gac	gtg	att	gtt	gtg	gac	tca	att	gag	tcc	gct	cga	3120
																Arg	
15	102					1030					1035					1040	
					-					٠							
20	aaa	ctg	cga	tca	atc	gat	gag	ttc	aag	tat	att	cca	att	gtt	ctc	tta	3168
				•								•				Leu	
					1045					1050		-			1055		
25																	
	gct	ccc	gtt	att	cat	gtc	agc	tta	aag	tct	gct	ttg	gat	ctt	ggt	atc	3216
30				•	His					-							
				1060	•				1065			÷		1070	-•		
35 .	act	tct	tac	atg	acc	act	cca	tgt	tta	acg	atc	gat	ctt	PRC.	aat	ggt	3264
					Thr												
40			1075					1080					1085	•			
																•	
	atg	att	cct	gct	ttg	gag	aat	cga	gct	gca	ccc	tca	ttg	BCB	gac	aac	3312
<b>4</b> 5					Leu												
•		1090					095					100					
50						-											
	aca	аая	tec	tto	gac	att	ctc	tto	gc^	<b>722</b>	<b>09</b> +	920	ato	ato	22+	cas	3360
					Asp												
55	~	<b></b> ;	J-01		,wh	× T C	LU	ren	UTG	GIM	nsp	Voll	114	191	VOII	ATII	

5	1105		1110		1	115	1120	
	cgc tt	a gcg gtg	aag att	cta gaa	aag tat	cac cac gtc	gtc aca gtc	3408
10	Arg Le	u Ala Val	Lys Ile	Leu Glu	Lys Tyr	His His Val	Val Thr Val	
			1125		1130		1135	
				•	-			٠
15	gtt ggo	c aat ggt	caa gaa	gca cta	gat gct a	atc aag gag	aaa cga tac	3456
	Val Gly	Asn Gly	Gln Glu	Ala Leu	Asp Ala	Ile Lys Glu	Lys Arg Tyr	
20		1140			145		1150	
							•	
	gat gtt	att ctc	atg gac	gtt cas	atg cca a	att atg gga	gga ttc gaa	3504
25							Gly Phe Glu	0001
		1155		1160		1165	ory The Ord	
30			•			1100		
	FC2 8C0	rct aag	att aga	g2g +2A	<b>739 077 0</b>	ort ott orn	acg caa aga	2550
	•						-	3552
35	- 1170				oin wig s	Ser Leu Gly	inr Gin Arg	
	. 1170	'	1	175		1180		
40	•							
40		•	•			tg ttg ggt		3600
	Thr Pro	Ile Ile	Ala Leu	Thr Ala	His Ala M	let Leu Gly	Asp Arg Glu	
45	1185	٠	1190		11	95	1200	
			• •	•	, t			
	aaa tgt	att caa	gcc caa a	atg gat	gaa tat c	tt tct aag	cct ctg aaa	3648
50	Lys Cys	Ile Gln	Ala Gln N	det Asp (	Glu Tyr L	eu Ser Lys	Pro Leu Lys	
		1	205		1210	•	1215	

	caa	aat	cat	ctt	att	cag	acg	atc	ttg	aaa	tgt	gca	acc	ctt	gga	ggt	3696
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				1220				•	1225					1230			
10																,	
	gça	ttg	ctc	gag	aag	ggt	agg	gag	gtt	agg	caa	tcc	gct	aat	gaa	gag	3744
											Gln						
15			1235					1240					1245				
										-							
20	agc	ccc	aat	tcg	caa	aat	ggt	cct	cgc	ggt	aca	cag	cat	cct	gca	tca	3792
											Thr						
		1250					1255					1260					
25	-				•												
	agt	ccc	aca	cca	gcc	cat	ato	а <i>та</i>	CCF	gct	atc	gaā	cct	cøt	ora.	tar.	3840
30											Ile						0010
30			•••	•••			IIIC C	, rr. 6				Olu	110	шБ		-	
	126	0	•			1270	•			•	1275				1	1280	
35															·	•	
-	acg	acc	act	ggc	cct	ata	aat	cat	gga	agt	gca	gag	agt	cct	tca.	ctt	3888
	Thr	Thr	Thr	Gly	Pro	Ile	Asn	His	Gly	Ser	Ala	Glu	Ser	Pro	Ser	Leu	
40				1	285				1	290				1	1295		
		٠		•													. •
45	gta	acg	gca	gat	gct	gag	gat	cca	ctt	gcg	agg	ctt	cta	atg	cgt	<b>g</b> cg	3936
	Val	Thr	Ala	Asp	Ala	G1u	Asp	Pro	Leu	Ala	Arg	Leu	Leu	Met	Arg	Ala	
			1	300				1	305				1	310			
50	•																
	cat	agc	agc	tag									÷				3948
	His	Ser	Ser											•	,		
55																	

1	વ	1	¢
1	v	4	i

•		
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•		
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	<213> Magnapotrthe grisea	
	water meditabout the Strong	
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	Ile Ala Thr Asn Ser Gly Ala Pro Gly Lys Asn Ala Ser Phe Arg Ser	
	20 25 30	

	Ser	Thr	Tyr	Val	Gln	Leu	Pro	Gly	Pro	G1u	Ser	Asp	Glu	Lys	Lys	Gln
5			35					40					45			
	Leu	Glu	Arg	Glu	Leu	Ala	Ala	Leu	Val	Ile	Arg	Val	Gln	GIn	Leu	G1u
10		50					55					60				
	Thr	Arg	Ala	Asn	Ala	A1a	Pro	Ala	Thr	Ile	Phe	Pro	Asp	Thr	Pro	Asn
45	65					70					75					80
15	Glu	Thr	Ala	His	Ser	Leu	Phe	G1 y	Asp	Asp	Ser	Ser	Ser	Pro	Thr	Ser
					85					90					95	
20	Ser	Ser	Ser	G1y	Arg	Glu	Pro	Lys	Arg	Leu	Lys	Ser	Ala	Ser	Ser	Thr
		-		100	•				105					110		
25	Thr	Arg		Gly	Phe	Thr	Thr	Asp	G1y	Arg	Pro	Ser	Lys	Leu	Asn	Ala
			115					120					125	•	•.	
	Ile	Thr	Asp	Glu	Glu	Leu	Glu	Gly	Leu	Arg	Glu	His	Val	Asp	Gly	Gln
30		130					135					140				
-		Arg	Leu	Leu	Asp	Ser	Gln	Arg	Ala	Glu	Leu	Asp	Gly	Val	Asn	Ala
 35	145	_				150					155					160
	Gln	Leu	Leu	Glu		Lys	Gln	Leu	G1n		Arg	Ala	Leu	Ala		Ile
	61				165			_		170		_	_		175	
40	Glu	Gln	Glu		Val	Ala	Thr	Leu		Arg	Glu	Leu	Trp		His	Gln
				180					185					190		
45	Lys	Ala		Glu	Ala	Phe	Gln		Ala	Leu	Arg	Glu		Gly	Ser	Ile
	V- 1	TVL	195		. 1		01	200		•			205	_	•.	
	Val		Ala	Ala	Ala			Asp	Leu	Ser	Lys	Arg	val	Lys	TIE	Asn
50	D	210	<b>~</b> 1	<b>M</b> . 4	A .		215		m)	æ)	D.	220		<b></b>		
		116	GIU	Met			GIU	TIE	ihr	lhr		Lys	Arg	Ihr		
55	225					230					235					240

	Ale	Met	Met	Asp	Gln	Leu	Gly	Val	Phe	Ser	Ser	Glu	Val	Ser	Arg	Yal
5					245					250					255	-
	Ala	Arg	Glu	Val	Gly	Thr	G1u	Gly	Ile	Leu	G1y	Gly	Gln	Ala	Gln	Ile
10				260					265					270		
	Glu	Gly	Val	Asp	Gly	Thr	Trp	Lys	G1u	Leu	Thr	Asp	Asn	Val	Àsn	Val
			275	-				280					285			
15	Met	Ala	Gln	Asn	Leu	Thr	Asp	Gln	Val	Arg	Glu	Ile	Ala	Ser	Val	Thr
		290					295					300				
20	Thr	Ala	Val	Ala	His	Gly	Asp	Leu	Thr	G1n	Lys	Ile	Glu	Ser	Ala	Ala
	305				•	310					315					320
25	Lys	Gly	G1u	Ile	Leu	Gln	Leu	Gln	Gln	Thr	Ile	Asn	Thr	Met	Val	Asp
25					325					330			-		335	
	Gln	Leu	Arg	Thr	Phe	Ala	Ser	Glu	Val	Thr	Arg	Val	Ala	Arg	Asp	Val
30				340					345					350		
	Gly	Thr	Glu	Gly	Met	Leu	G1y	Gly	Gln	Ala	Asp	Val	Glu	Gly	Val	Lys
35 .		•	355					360					365			
-	Gly	Met	Trp	Asn	Glu	Leu	Thr	Val	Asn	Val	Asn	Ala	Met	Ala	Asn	Asn
		370					375					380				
40	Leu	Thr	Thr	G1n	Val	Arg	Asp	Ile	Ile	Asn	Val	Thr	Thr	Ala	Val	Ala
	385					390					395					400
45	Lys	Gly	Asp	Leu	Thr	G1n	Lys	Val	Gln	Ala	G1u	Cys	Arg	Gly	Glu	Ile
					405					410			•		415	•
	Phe	Glu	Leu	Lys	Asn	Thr	Ile	Asp	Ser	Met	Val	Asp	G1n	Leu	Gln	Gln
50				420					425					<b>43</b> 0		
	Phe	Ala	Arg	G1u	Val	Thr	Lys	Ile	Ala	Arg	Glu	Val	Gly	Thr	Glu	Gly
55		•	435					440			-		445			

	Arg	Leu	Gly	Gly	Gln	Ala	Thr	Val	His	Asp	۷al	Gln	Gly	Thr	Trp	Arg
5		450					455					460				
	Asp	Leu	Thr	Glu	Asn	Yal	Asn	G1y	Met	Ala	Met	Asn	Leu	Thr	Thr	Gln
10	465					470					475					480
	Val	Arg	Glu	Ile	Ala	Asn	Val	Thr	Ser	Ala	Val	Ala	Ala	Gly	Asp	Leu
					485					490			•		495	
15	Ser	Lys	Lys	Ile.	Arg	Val	Glu	Val	Lys	Gly	Glu	Ile	Leu	Asp	Leu	Lys
				500			•		505					510		
20	Asn	Thr	Ile	Asn	Thr	Met	Yal	Asp	Arg	Leu	Ġly	Thr	Phe	Ala	Phe	Glu
•			515					520	•				525			
25	Val	Ser	Lys	Val	Ala	Arg	Ala	Val	G1y	Thr	Asp	Gly	Thr	Leu	Gly	G1y
23		530					535					540				
	Gln	Ala	Gln	Val	Glu	Asn	Val	G1u	Gly	Lys	Trp	Lys	Asp	Leu	Thr	G1u
30	545					550					555					560
•	Asn	Va1	Asn	Thr	Met	Ala	Ser	Asn	Leu	Thr	Ser	Gln	Val	Arg	Gly	Ile
35					565					570					575	
-	Ser	Thr	Val	Thr	Gln	Ala	Ile	Ala	Asn	Gly	Asp	Met	Ser	Arg	Lys	Ile
-				580					585					<b>59</b> 0		
40	Asp	Val		Ala	Lys	Gly	Glu	Ile	Leu	Ile	Leu	Lys	Glu	Thr	Ile	Asn
			595					600	-				605			
45	Asn		Val	Asp	Arg	Leu	Ser	Ile	Phe	Cys	Asn	Glu	Val	Gln	Arg	Val
		610					615					620				
		Lys	Asp	Val			Asp	Gly	Ile	Met		Gly	Gln	Ala	Asp	Val
50	625			,		630					635					640
	Ala	Gly	Leu			Arg	Trp	Lys	Glu		Thr	Thr	Asp	Val	Asn	Thr
55					645					650					<b>65</b> 5	

	Met	Ala	Asn	Asn	Leu	Thr	Ala	Gln	Val	Arg	Ala	Phe	G1 y	Asp	Ile	Thr
5		•		660					665					670		
	Asn	Ala	Ala	Thr	Asp	Gly	Asp	Phe	Thr	Lys	Leu	Val	G1u	Val	Glu	Ala
10			675					680					<b>68</b> 5			
	Ser	Gly	Glu	Met	Asp	Glu	Leu	Lys	Arg	Lys	Ile	Asn	Gln	Met	Val	Tyr
		690					695					700				
15	Asn	Leu	Arg	Asp	Ser	Ile	Gln	Arg	Asn	Thr	Gln	Ala	Arg	Gĺu	Ala	Ala
	705		-			710					715					720
20	Glu	Leu	Ala	Asn	Lys	Thr	Lys	Ser	Glu	Phe	Leu	Ala	Asn	Met	Ser	His
				•	725					730			•	•	735	
25	Glu	Ile	Arg	Thr	Pro	Met	Asn	Gly	Ile	Ile	Gly	Met	Thr	Gln	Leu	Thr
25				740					745					750		
	Leu	Asp	Thr	Asp	Leu	Thr	Gln	Tyr	G1n	Arg	Glu	Met	Leu	Asn	Ile	Val
30			<b>75</b> 5					760					765			
	Asn	Asn	Leu	Ala	Met	Ser	Leu	Leu	Thr	Ile	Ile	Asp	Asp	Ile	Leu	Asp
 35		770				•	7 <b>7</b> 5					780	•			
-	Leu	Ser	Lys	Ile	Glu	Ala	Lys	Arg	Met	Val	Ile	Glu	Glu	Ile	Pro	Tyr
	785					790					795					800
40	Thr	Leu	Arg	.G1y	Thr.	Val	Phe	Asn	Ala	Leu	Lys	Thr	Leu	Ala		Lys
					805					810			-		815	
45	Ala	Asn	Asp		Phe	Leu	Asp	Leu		Tyr	Arg	Val	Asp		Ser	Va1
				820					825	-				830		
	Pro	Asp <sub>.</sub>	His	Val	Ile	Gly	Asp	Ser	Phe	Arg	Leu	Arg	Gln	Ile	Ile	Leu
50			835					840					845			
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		Leu	Thr	Ile	Gln	Lys	Gly	Asn	Asp	Val	Thr	Cys	Leu	Pro	Asn	Glu	Tyr
5		865					870					875					880
		Met	Ile	G1u	Phe	Va1	Val	Ser	Asp	Thr	Gly	Ile	G1y	Ile	Pro	Thr	Asp
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		Lys	Leu	Gly	Leu	Ile	Phe	Asp	Thr	Phe	G1n	Gln	Ala	Asp	Gly	Ser	Met
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				915					920					925		•	
. 20		Leu	Val	Asn	Leu	Met	Gly	Gly	Asp	Val	Trp	Val	Lys	Ser	Gln	Tyr	Gly
			930					935					940				
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25		945					950					955					960
		Asp	Ile	Ser.	Leu	Ile	Arg	Lys	Gln	Leu	Lys	Pro	Tyr	Lys	Gly	His	Gln
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		Val	Leu	Phe	Ile	Asp	Lys	Gly	Lys	Thr	Gly	His	Gly	Pro	Glu	Val	Gly
35					980					985	-				990		
-		Gln	Met	Leu	Gly	Gln	Leu	G1y	Leu	Val	Pro	Ile	Val	Leu	Glu	Ser	Glu
				995				:	1000				1	1005			
40		Gln	Asn	His	Thr	Leu	Thr	Arg	Val	Arg	Gly	Lys	Glu	Cys	Pro	Tyr	Asp
		1	1010				1	1015				1	1020				
45		Val	Ile	Val	Val	Asp	Ser	Ile	Asp	Thr	Ala	Arg	Arg	Leu	Arg	Gly	Ile
	٠	1028	5	•		1	1030		٠		]	1035				1	L <b>04</b> 0
		Asp	Asp	Phe	Lys	Tyr	Leu	Pro	Ile	Val	Leu	Leu	Ala	Pro	Thr	Val	His
50					1	045		•		1	1050				1	LO55	
		Val	Ser	Leu	Lys	Ser	Cys	Leu	Asp	Leu	Gly	Ile	Thr	Ser	Tyr	Met	Thr
55				. ]	1060				1	065				;	1070		

	Met Pro	Cys Lys	Leu Ile	Asp Leu	Gly Asn	Gly Met	Val Pro Ala Leu
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	Glu Asn	Arg Ala	Thr Pro	Ser Leu	Ser Asp	Asn Thr	Lys Ser Phe Glu
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	Ile Leu	Leu Ala	Glu Asp	Asn Thr	Val Asn	Gln Arg	Leu Ala Val Lys
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15	Ile Leu	Glu Lys	Tyr Asn	His Val	Val Thr	Val Val	Ser Asn Gly Ala
		:	1125	·	1130		1135
20	Glu Ala	Leu Glu	Ala Val	Lys Asp	Asn Lys	Tyr Asp	Val Ile Leu Met
	•	1140			1145		1150
25	Asp Val	Gln Met	Pro Val	Met Gly	Gly Phe	Glu Ala	Thr Ala Lys Ile
25	1	1155		1160		:	1165
	Arg Glu	Tyr Glu	Arg Ser	Leu Gly	Thr Gln	Arg Thr	Pro Ile Ile Ala
30	1170		. 1	175		1180	
. •	Leu Thr	Ala His	Ala Met	Met Gly	Asp Arg	Glu Lys	Cys Ile Glu Ala
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-	Gln Met	Asp Glu	Tyr Leu	Ser Lys	Pro Leu	Gln Gln	Asn His Leu Ile
		. 1	205		1210		1215
40	Gln Thr	Ile Leu	Lys Cys	Ala Thr	Leu Gly	Gly Ala	Leu Leu Glu Gln
		1220		. 1	.225		1230
45	Asn Arg	Glu Arg	Glu Leu	Glu Leu	Ala Arg	His Ala	Glu His Lys Gly
	1	235		1240		1	.245
	Gly Leu	Ser Thr	Asp Pro	Ala Arg	Ala Ser	Ser Val	Met Arg Pro Pro
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	Leu His	His Arg	Pro Val	Thr Thr	Ala Glu	Ser Leu	Ser Gly Gly Ala
55	1265		1270		1	275	1280

Glu Ser Pro Ser Leu Met Ala Asn Asp Gly Glu Asp Pro Ile Gln Arg

Ala Arg Ser Ser Leu Ser Glu Pro Gly Cys Leu <210> 17 <211> 3924 <212> DNA <213> Magnapotrthe grisea <220> <221> CDS <222> (1).. (3924) <400> 17 atg gcg gac gcg gcg act ctg gca gct gtc gct gcg att gtg gag aat Met Ala Asp Ala Ala Thr Leu Ala Ala Val Ala Ala Ile Val Glu Asn atc gct acc aac tcg ggg gcc cct gga aaa aat gct tca ttt cgc tcc Ile Ala Thr Asn Ser Gly Ala Pro Gly Lys Asn Ala Ser Phe Arg Ser agt acc tat gtc cag ctt ccc ggt ccg gaa tcc gac gag aag aaa cag Ser Thr Tyr Val Glm Leu Pro Gly Pro Glu Ser Asp Glu Lys Lys Gln 

			35					40					45	•			
5																	
	ctc	gag	cgc	gag	ctt	gcc	gcc	ctg	gtg	ata	agg	gta	cag	cag	ctc	gaa	192
10	Leu	Glu	Arg	Glu	Leu	Ala	Ala	Leu	Val	Ile	Arg	Val	Gln	Gln	Leu	Glu	
		50					55					60					
15	acc	cgt	gcc	aac	gcg	gct	cct	gct	aca	ata	ttc	ccc	gac	aca	ccc	aac	240
	Thr	Arg	Ala	Asn	Ala	Ala	Pro	Ala	Thr	Ile	Phe	Pro	Asp	Thr	Pro	Asn	
20	65					70					75					80	•
								٠.									
	gaa	act	gca	cat	tca	ctc	ttt	ggc	gat	gat	agc	tcg	tcc	cct	acc	agt	288
25	Glu	Thr	Ala	His	Ser	Leu	Phe	Gly	Asp	Asp	Ser	Ser	Ser	Pro	Thr	Ser	
					85				÷	90	•				95		
30			•														
-	tcg	agc	tca	ggc	cgg	gag	cct	aaa	cga	ctg	aag	tcg	gca	tcc	agc	aca	336
	Ser	Ser	Ser	Gly	Arg	Glu	Pro	Lys	Arg	Leu	Lys	Ser	Ala	Ser	Ser	Thr	
35 ·				100				•	105					110			
•																	
40	acg	agg	aat	ggt	ttc	act	acg	gac	ggt	cgt	cca	tca	8ag	ctc	aac	gca	384
	Thr	Arg	Asn	Gly	Phe	Thr	Thr	Asp	Gly	Arg	Pro	Ser	Lys	Leu	Asn	Ala	
	٠		115					120			-		125				
45			•														٠
	atc	acc	gat	gag	gag	ctc	gaa	ggc	ttg	cgc	gaa	cat	gtt	gac	ggc	cag	432
50	Ile	Thr	Asp	Glu	Glu	Leu	Glu	Gly	Leu	Arg	G1u	His	Val	Asp	Gly	Gln	
		130					135					140					

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10																	
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	G1n	Leu	Leu	Glu	Gln	Lys	Gln	Leu	Gln	Glu	Arg	Ala	Leu	Ala	Ile	Ile	
15	•				165					170					175		
					٠.												
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	G1u	Gln	Glu	Arg	Val	Ala	Thr	Leu	G1u	Arg	Glu	Leu	Trp	Lys	Hïs	Gln	
25				180					185					190			
?5														•			
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30	Lys	Ala	Asn	Glu	Ala	Phe	Gln	Lys	Ala	Leu	Arg	G1u	Ile	Gly	Ser	Ile	
			195					200					205				
35															_	÷	
-	gtg	acc	gct	gca	gcc	cgg	ggt	gac	ctc	tct	aag	agg	gtc	aag	ata	вас	672
	Va1	Thr	Ala	Ala	Ala	Arg	Gly	Asp	Leu	Ser	Lys	Arg	Va1	Lys	Ile	Asn	
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										•							
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	Pro	Ile	Glu	Met	Asp	Pro	G1u	Ile	Thr	Thr	Phe	Lys	Arg	Thr	Met	Asn	
	225					230					235				•	240	
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5	Ala	Met	Met	Asp	G1n	Leu	Gly	Val-	Phe	Ser	Ser	Glu	Val	Ser	Arg	Val	

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10						٠.				cag Gln	atc Ile	816
15										aac Asn		864
20			275			280			285	ŧ	•	
25										gtc Val		912
30	000	go+		 	 	<b>*</b> * -				 		060
35 -	_					_	•			gcg		960
40				Leu						gtg Val		1008
45				325			330			335		
50										gac Asp		1056

5	gga	acc	gag	gga	atg	ctc	ggc	ggg	cag	gct	gac	gtt	gaa	ggg	gtc	aag	1104
	Gly	Thr	Glu	Gly	Met	Leu	Gly	Gly	Gln	Ala	Asp	Val	Glu	Gly	Val	Lys	
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10																	
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	Gly	Met	Trp	Asn	Glu	Leu	Thr	Val	Asn	Val	Asn	Ala	Met	Ala	Asn	Asn	
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	Leu	Thr	Thr	Gln	Val	Arg	Asp	Ile	Ile	Asn	Val	Thr	Thr	Ala	Val	Ala	
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			•														
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35			•														
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	Phe	G1u	Leu	Lys	Asn	Thr	Ile	Asn	Ser	Met	Val	Asp	G1n	Leu	Gln	Gln	
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	Phe	Ala	Arg	Glu	Val	Thr	Lys	Ile	Ala	Arg	Glu	Val	G1y	Thr	Glu	Gly	
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	465					470					475					480	
		•															
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•	Val	Arg	G1u	Ile	Ala	Asn	Val	Thr	Ser	Ala	Val	Ala	Ala	Gly	Asp	Leu	
20				•	485					490					495		
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25	Ser	Lys	Lys	Ile	Arg	Val	Glu	Val	Lys	G1y	G1u	Ile	Leu	Asp	Leu	Lys	
			•	500					505			٠		510			
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•	Val	Ser	Lys	Va1	Ala	Arg	Ala	Val	Gly	Thr	Asp	Cly	Thr	Leu	Ġly	Gly	
		530					535					540					
45																	
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50	Gln	Ala	Gln	۷al	Glu	Asn	Val	G1u	G1y	Lys	Trp	Lys	Asp	Leu	Thr	Glu	-
	545		•			550					555					560	·

	aa	c gt	c aac	aco	e atg	gcg	tca	aac	cto	act	tct	cag	gto	ag	g gga	a ata	1728
5	Ası	n Val	l Asr	1 Thu	Met	Ala	Ser	Asn	Lei	ı Thr	Ser	Glr	(Va)	Ar	g G13	rÎle	
					565	i				570	)				578	5	
10																	•
	tea	a acc	gtg	aca	caa	gcc	atc	gcg	aac	ggt	gac	ate	ago	cga	aag	atc	1776
					Gln												
15				580	-				585					590			
					•												
20 .	gao	gtg	gaa	gcc	aag	ggc	gag	ata	cta	atc	ctc	aag	gaa	act	atc	aac	1824
	Asp	Val	G1u	Ala	Lys	Gly	Glu	Ile	Leu	Ile	Leu	Lys	Glu	Thr	·Ile	Asn	
			595					600					605				
25																	
	aac	atg	gtt	gat	cgt	ctg	tcg	ata	ttc	tgc	aat	gaa	gta	caa	cga	gtc	1872
30					Arg	•											
		610					615					620			_		
										•			·				
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					Gly												
10	625				•	630	•	•			635			0	,,,,,	640	
								٠								0.10	
	gca	ggt	ctc	aag	ggg	CFA	tao	226	ga g	att	acc	200	oat.	at c	920	200	1968
<b>15</b> .					Gly								•				
•		,		2,0	645		p	<i>D</i> , 3	<b>J10</b>	650	110.	1111	vaħ	141	655	LITT	
50					040			•		050	-				000		
	2 🕇 🕾	<b>70</b> 0	026	201	.++				_+_			**-					0010
					ctt											-	2016
5	me t	LIA	กรม	usii	Leu	III	ита	GIU	val	HIE	W18	rne	GTA	Asp	TIG	inr	

aat gcc gct acc gac gga gac ttc acc aag ctg gtc gag gtt gag gcg Asn Ala Ala Thr Asp Gly Asp Phe Thr Lys Leu Val Glu Val Glu Ala tog ggc gaa atg gac gaa ctg aag cgc aag atc aat caa atg gtc tac Ser Gly Glu Met Asp Glu Leu Lys Arg Lys Ile Asn Gln Met Val Tyr aat ctc cga gac agt atc caa aga aac acg caa gca aga gaa gcc gca Asn Leu Arg Asp Ser Ile Gln Arg Asn Thr Gln Ala Arg Glu Ala Ala gas ttg gcc aac aag acg aag tcg gag ttc ctc gct aac atg tcc cac Glu Leu Ala Asn Lys Thr Lys Ser Glu Phe Leu Ala Asn Met Ser His gaa atc cgc aca ccc atg aac ggt atc atc ggc atg aca caa ctt act Glu Ile Arg Thr Pro Met Asn Gly Ile Ile Gly Met Thr Gln Leu Thr tt gat aca gat ttg acg caa tac caa cgc gaa atg ctc aac att gtc Leu Asp Thr Asp Leu Thr Gln Tyr Gln Arg Glu Met Leu Asn Ile Val 760 . 

E	aac	aat	cto	gcc	atg	agt	ctg	cto	acc	att	atc	gac	gac	atc	ctc	gat	2352
5	Asn	Asn	Leu	Ala	Met	Ser	Leu	Leu	Thr	Ile	Ile	Asp	Asp	Ile	Leu	Asp	
		770	١				775					780					
10																	
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4.5	Leu	Ser	Lys	Ile	G1u	Ala	Lys	Arg	Met	Val	Ile	Glu	Glu	Ile	Pro	Tyr	
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20	acg	tta	cga	gga	acg	gtc	ttc	aac	gca	ctg	aag	act	ttg	gcg	gtc	aag	2448
	Thr	Leu	Arg	G1y	Thr	Val	Phe	Asn	Ala	Leu	Lys	Thr	Leu	Ala	Val	Lys	
25					805					810		•			815		
25															. *		
	gcg	aac	gac	aag	ttt	ttg	gat	ctc	acg	tac	cgt	gtg	gac	agc	tca	gtt	2496
30	Ala	Asn	Asp	Lys	Phe	Leu	Asp	Leu	Thr	Tyr	Arg	Val	Asp	Ser	Ser	Val	
-				820		•			825					830			
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	Pro	Asp	His	Val	Ile	Gly	Asp	Ser	Phe.	Arg	Leu	Arg	Gln	Ile	Ile	Leu	
40			835		-			840					845				
													•				
45	aac	ctg	ġtt	ggc	aat	gcc	atc	a8a	ttc	acc	gag	cat	gga	gag	gtc	agc	2592
									Phe								
		850					<b>85</b> 5					860	-				
50																	
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<i>EE</i>												•				u Tyr	
55					-						•						

5	865					870					875					880	
10					Val					Gly	ata Ile				Thr	_	2688
15	222	ctg	ggt	ctc	885 atc	ttc	gaç	aca	ttc	890 cag	cag	gct	gat	gga	895 tcc	atg	2 <b>73</b> 6
<b>20</b>	Lys	Leu	Gly	Leu 900	Ile	Phe	Asp	Thr	Phe 905	Gln	Gln	Ala	Asp	Gly 910	Ser	Met	
25			Lys					Gly ·			ctg Leu		Ile				2784
30	ctc	gtc	915 aac	ctc	atg	ggc	ggt	920 gac	gtg	tgg	gtc	aag	925 tca	caa	tac	ggc	2832
35 -	Leu	Val 930	Asn	Leu	Met	Gly	Gly 935	Asp	Val	Trp	Val	Lys 940	Ser	Gln	Тут	Gly	
40							•				gtc Val		•	•			2880
45	945			•		950					955					960	
50											cct Pro			Gly			2928

	gto	ct	g tt	c ato	gat	aag	ggo	888	g act	gg:	a cad	ggg	g cc	c ga	g gt	g ggg	2976
5	Va]	Le	ı Ph	e Ile	Asp	Lys	Gly	Lys	Thi	G13	/ His	G13	Pre	o Gl	u Va	l Gly	
				980	)				985	5				99	0		
10																	
	cag	ate	cto	ggc	cag	ctg	ggt	ttg	gtg	ccc	ato	gte	cte	gaa	a tc	gag	3024
	Gln	Met	: Le	ı Gly	Gln	Leu	Gly	Leu	Val	Pro	Ile	Val	Lei	1 G1	ı Sei	r Glu	
15				99	5				100	0 .				100	)5		
<b>20</b>	caa	aat	cac	acc	ctg	acg	cgg	gtg	cgc	ggc	aag	gaa	tgt	ccc	tac	gac	3072
	Gl'n	Asn	His	Thr	Leu	Thr	Arg	Val	Arg	Gly	Lys	Glu	Cys	Pro	Тут	Asp	
		1010					1015					1020					
25																	
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30	Val	Ile	Val	Val	Asp	Ser	Ile	Asp	Thr	Ala	Arg	Arg	Leu	Arg	Gly	Ile	
	102	5			;	1030				:	1035					1040	
	•																
<b>35</b> .	gac	gac	ttc	aag	tat	ctg	ccc	atc	gtt	ctc	ctg	gcg	cca	act	gtc	cac	3168
	Asp	Asp	Phe	Lys	Tyr	Leu	Pro	Ile	Val	Leu	Leu	Ala	Pro	Thr	Val	His	
40				1	1045				1	1050					1055		
																	•
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45	Val	Ser	Leu	Lys	Ser	Cys	Leu	Asp	Leu	Gly	Ile	Thr	Ser	Tyr	Met	Thr	
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50									٠								
	atg	CCC	tgc	aag	ctc	atc	gac	ctc	ggc	aat	ggt	atg	gtt	ccc	gct	ctt	3264
	Met	Pro	Cys	Lys	Leu	Ile	Asp	Leu	Gly	Asn	Gly	Met	Val	Pro	Ala	Leu	
55																	

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10	Glu Asn	Arg Ala	Thr Pro S	er Leu Ser	Asp Asn Thr Lys S	er Phe Glu
	1090		. 10	95	1100	٠.
15					aac cag cgc ctg g	-
		Leu Ala	Glu Asp A	sn Thr Val	Asn Gln Arg Leu A	la Val Lys
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25	lle Leu			is Val Val	Thr Val Val Ser A	sn Gly Ala
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30						
-					aaa tac gat gtg a	
	Glu Ala		Ala Val Ly		Lys Tyr Asp Val I	le Leu Met
35		1140		1145	11	50
40	•				ttt gag gcg acg g	_
			Pro Val Me		Phe Glu Ala Thr A	la Lys Ile
45	,	155		1160	1165	·
73			:			2550
					cag agg aca cca at	
50		iyr Giu			Gln Arg Thr Pro II	le lle Ala
	1170		117	, d	1180	

	ctt	aco	gct	cac	gca	atg	atg	ggc	gao	cgt	gag	; aag	tgt	ato	gaı	g gcc	3600
5	Leu	Thr	Ala	His	Ala	Met	Met	Gly	Asp	Arg	Glu	Lys	Cys	Ile	G1	ı Ala	
	118	5				1190					1195	i				1200	
10																	
,,	cag	ate	gac	gag	tac	ctg	tcg	aag	cct	ctg	cag	cag	aac	cac	ttg	ata	3648
	Gln	Met	Asp	G1u	Tyr	Leu	Ser	Lys	Pro	Leu	Gln	Gln	Asn	His	Leu	lle	
15					1205					1210					1215	;	
													-				
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	Gln	Thr	Ile	Leu	Lys	Cys	Ala	Thr	Leu	G1 y	G1y	Ala	Leu	Leu	Glu	Gln	
				1220	•				1225				:	1230			
25			٠	-									٠				
	aat	cgt	gag	cgc	gag	ctt	gas	cta	gca	agg	cat	gcc	gaa	cac	aaa	gga	3744
30	Asn	Arg	Glu	Arg	Glu	Leu	Glu	Leu	Ala	Arg	His	Ala	Glu	His	Lys	Gly	
			1235					1240					1245				
35																cca	37 <del>9</del> 2
			Ser	Thr	Asp		•	Arg	Ala	Ser	Ser	Val	Met	Arg	Pro	Pro	
40		1250				. ]	1255				1	1260					•
45					ccg												3840
45			His	Arg	Pro		Thr	Thr	Ala			Leu	Ser	Gly			
	1265	•			1	270				]	1275					1280	
50																	
					ttg												3888
55	GIU	ser	rro	ser	Leu	Met	Ala	Asn	Asp	Gly	Glu	Asp	rro	116	GIN	Arg	

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15	Z010\ 00	
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	Asp	Thr	Leu	Phe	Gly	Glu	Glu	Ala	Gln	Ala	Val	Ala	Val	Arg	Pro	Lys
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3 <i>0</i>				100					105					110		
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	545					550					555					560
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5	Ser	· G1r	ı Val	Glr	Let	ı Pro	G13	Pro	Asg	Th	Pro	Ala	Lys	s Arg	g Lys	s Leu	
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10							•				•						
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		Ala	Ser	Ala	Thr	Ser	Pro	Phe	Pro	Glu	Thr	Pro	Asn	Glu	Val	Ile	
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-											cac						336
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	Ser	Tyr	Gln	Leu	Thr	Glu	Glu	Ala	Leu	Glu	Gly	Leu	Arg	Glu	His	Val	
			115					120					125				· •
io																	
											cgc						432
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45	Arg	Met 210	Asn	Thr	Val	Glu	Met 215	Asp	Pro	Glu	Ile	Thr 220	Thr	Phe	Lys	Arg	
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	225	114	u9II	ut <b>q</b>		met 230	ush	O1II	TAN		235	- MS	VTS	⊃€Ľ	<b>A</b> III	va1 240	

	tcg c	ga gt	c gct	cgt	gaa	gto	gg1	t acc	gaa	a gga	tte	ctt	ggt	gg	caa	768
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				245	i				250	)				258	5	
10																
	gcc c	gt ato	ggc	ggc	gto	gac	gga	aca	tgg	aag	gaa	ttg	act	gac	aac	816
	Ala Ai	rg Ile	• Gly	Gly	Val	Asp	Gly	Thr	Trp	Lys	Glu	Leu	Thr	Asp	Asn	
15			260					265					270			
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	Val As			-												
		275					280				-	285				
25							•				•					
	tcg gt	t acc	acc	gcc	gtg	gcc	cac	ggc	gat	ctg	act	823	aag	atc	gaa	912
30	Ser Va	l Thr	Thr	Ala	Val	Ala	His	G1y	Asp	Leu	Thr	Lys	Lys	Ile	G1u	
	29	0 .	•	•		295					300				-	
-																
<b>35</b> -	cga cc	t gcc	aga	ggc	gag	ata	ttg	caa	tta	caa	сва	acg	att	aac	acc	960
	Arg Pr	o Ala	Arg	G1y	Glu	Ile	Leu	Gln	Leu	Gln	G1n	Thr	Ile	Asn	Thr	
40	305				310					315					320	
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C.	atg gt	g gac	caa	tta	cga	aça	ttt	gct	tct	gaa	gtc	aca	cgt	gta	gcg	1008
<b>45</b> .	Met Va	l Asp	G1n	Leu	Arg	Thr	Phe	Ala	Ser	Glu	Val	Thr	Arg	Val	Ala	
	•			325					330					335		
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	aga gat	gtc	ggg	acc .	gaa	ggc	atg	tta	ggc	ggg	caa	gcc .	gat	gtt	ggg	1056
	Arg Asp															
55																

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	Ala	Asn	Asn	Leu	Thr	Thr	Gln	Val	Arg	Asp	Ile	Ile	Lys	Val	Thr	Thr	
20		370	•				375					380		• •			•
									÷								
	gct	gtc	ECC	aag	gga	gat	ctt	аса	сая	aag	gtc	caa	gcc	gat	tec	200	1200
25 .					_					_	_	Gln		_		•	
	385			2,0	<b>02</b> 7	390			0111	2,0	395	<b>0111</b>	1144.	uap	0,3	400	
	000					,					000			·		400	
30	aga	<i>σ</i> ο <i>σ</i>	ata	++0	<b>4</b> 0.4	a ta	226	***	000	a <b>t</b> 0	200	tcc	a+~			<b>4</b> 0 ÷	1940
-									•				_		_		1248
35	Gly	Old	TIE	rue		Leu	Lys	261	ш		ASII	Ser	Me C	val		GIN	
-					405					410					415		
												•					
40									-			gcc					1296
	Leu	Gln	Gln	Phe	Ala	Arg	Glu	Val	Thr	Lys	Ile	Ala	Arg	Glu	Val	Gly	
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	acc	tgg	agg	gat	ctg	8cg	gag	aac	gto	aac	ggc	atg	gcc	atg	aac	ttg	1392
5	Thr	Trp	Arg	Asp	Leu	Thr	G1u	Asn	Val	Asp	Gly	Met	Ala	Met	Asn	Leu	
		450					455					460					
10		•					٠					-					
	acc	act	çaa	gtg	cga	gaa	att	gcc	aag	gtt	aca	aca	gct	gtc	gcc	822	1440
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														•			
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25																	
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	Ala	Val	Glu	Val	Ser	Lys	Val	Ala	Arg	G1u	Val	Gly	Thr	Asp	Gly	Thr	
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	Leu	Gly	Gly	Gln	Ala	G1n	Val	Ala	Asn	Val	Glu	Gly	Lys	Trp	Lys	Asp	
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50																	
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55	Leu	Thr	Glu	Asn	Val .	Asn	Thr	Met	Ala	Ser	Asn	Leu	Thr	Val	G1n	Val	

5	545					550	•				555		٠.			560	
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30																	
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35 -		610					615				•	620	•				
•		•	•			•	-										•
40	gcc	gac	gta	ggt	ggt	cta	gac	ggc	cgc	tgg	aaa	gag	atc.	acc	aca	gat	1920
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Ω.	625					630					635					640	
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	Met	Ile	Ser	Asn	Leu	Arg	Asp	Ser	Ile	Gln	Arg	Asn	Thx	Gln	Ala	Arg	
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25																	
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30	Glu	Ala	Ala	Glu	Leu	Ala	Asn	Lys	Thr	Lys	Ser	Glu	Phe	Leu	Ala	Asn	
	705					710			-	-	715				•	720	
-	•																
35 -	atg	tcc	cat	gaa	att	cga	acg	ccg	atg	aac	ggt	atc	atc	gga	- atg	act	2208
	Met	Ser	His	Glu	Ile	Arg	Thr	Pro	Met	Asn	Gly	Ile	Ile	Gly	Met	Thr	
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50																•	
	aac	att	gtc	aat	aat	ctt	gcc	aat	agc	ctc	ttg	acg	ata	att	gac	gat	2304
								Asn									·
55															-	-	

5			755	5		٠.		760	ı				765				
	ato	ttg	gat	ctt	tcc	aag	att	gas	gct	cgg	aga	atg	gtc	att	gag	gag	2352
10	Ile	Leu	Asp	Leu	Ser	Lys	Ile	Glu	Ala	Arg	Arg	Met	Val	Ile	G1u	Glu	
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15							-										
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	Ile	Pro	Tyr	Thr	Leu	Arg	Gly	Thr	Val	Phe	Asn	Ala	Leu	Lys	Thr	Leu	
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25	gct	gtc	aag	gca	aat	gag	aag	ttc	ttg	gat	ctc	acc	tac	aag	gtc	gat	2448
	Ala	Val	Lys	Ala	Asn	Glu	Lys	Phe	Leu	Asp	Leu	Thr	Tyr	Lys	Val	Asp	
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30											*						
-	agc	tcc	gtg	cct	gac	tac	gtt	att	ggc	gac	tcc	ttc	cgt	ctc	aga	caa	2496
35	Ser	Ser	Val	Pro	Asp	Tyr	Val	Ile	Gly	Asp	Ser	Phe	Arg	Leu	Arg	Gln	
-				820					825	•				830			
40									•							•	
••				aac													2544
	He	Ile		Asn	Leu	Val	Gly	_	Ala	Ile	Lys	Phe	Thr	Glu	His	Gly	
45			835					840					845		•		
		8.7												•			4
50				cta													2592
			Ser	Leu	Thr			Glu	Ser	Met			Asn	Asn	Val	Arg	
		850					855					860	•				

		cct	gga	gag	tat	gcg	gtt	gag	ttt	gto	gtg	gag	gac	acg	ggc	ata	gga	2640
5		Pro	Gly	Glu	Tyr	Ala	Val	Glu	Phe	Val	Va1	Glu	Asp	Thr	Gly	Ile	Gly	
		865	;				870					875	;				880	
10		atc	gco	caa	gat	aaa	ctg	gat	ttg	atc	ttc	gac	acg	ttc	caa	саа	ØC Ø	2688
						Lys												2000
					p	885	200	щ	LUG	110	890			1110	0111		vra	
15						003					930					895		
20										٠.								2736
		Asp	Gly	Ser	Met	Thr	Arg	Lys	Phe	Gly	Gly	Thr	Gly	Leu	Gly	Leu	Ser	
					900					905					910			
25		٠															•	
		att	tcg	aaa	cga	ctc	gtc	aat	ctc	atg	ggt	ggt	gat	ctc	tgg	gta	aac	2784
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30				915					920					925				
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· 35		agt	gaa	cat	ggc	aag	ggc	agt	gaa	ttt	cac	ttc	aça	tgc	tta	gtc	aag	2832
	-	Ser	Glu	His	Gly	Lys	Gly	Ser	Glu	Phe	His	Phe	Thr	Cys	Leu	Val	Lys	
			930	•				935					940					
40							•									ė		
	•	ctt	get	cet	gac	gat	gct	gct.	cte	atc	σaσ	caa	CBB	atc	COC.		tac	2880
45						Asp												2000
		945	MIG	110	nsp			NIZ	Leu	116	Ulu	•	GIII	116	VT R			
		340				•	950					955				•	960	
50																		
						gtg												2928
		Arg	Gly	His	Gln	Val	Leu	Phe	Val	Asp	Lys	Ala	Gln	Ser	Gln	Asn	Ala	
55																		•

					965					970	•				975		
5											•						
	ace	e tea	ato	aag	cct	atg	ctt	gag	aag	atc	ggg	ctg	aag	cct	gtc	gtt	2976
	Thu	Ser	Ile	Lys	Pro	Met	Leu	Glu	Lys	Ile	Gly	Leu	Lys	Pro	Val	Val	
10				980					985					990			
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15	gte	gat	tcg	gag	aag	agt	cct	gcg	ctg	act	cgt	ctt	caa	agc	ggt	ggc	3024
	Va]	Asp	Ser	Glu	Lys	Ser	Pro	Ala	Leu	Thr	Arg	Leu	Gln	Ser	Gly	Gly	
			995					1000				;	1005				
20													·				
	tco	ctt	ccc	tat	gat	gct	atc	ctc	gtc	gat	tcc	atc	gac	act	gcg	aga	3072
25	Ser	Leu	Pro	Tyr	Asp	Ala	Ile	Leu	Va1	Asp	Ser	Ile	Asp	Thr	Ala	Arg	
		1010				1	1015					1020					
30	agg	tta	aga	gcc	gtg	gac	gat	ttc	aag	tac	ctt	cct	atc	gtc	ttg	ctg	3120
	Arg	Leu	Arg	Ala	Val	Asp	Asp	Phe	Lys	Tyr	Leu	Pro	Ile	Val	Leu	Leu	
- : 35	102	5			]	030				]	1035				1	1040	
	-																
	gca	cca	gtt	gtt	cac	gtt	agt	ctg	aag	tcg	tgc	ttg	gat	ctg	gga	att	3168
40	Ala	Pro	Val	Val	His	Val	Ser	Leu	Lys	Ser	Cys	Leu	Asp	Leu	Gly	I1e	
				1	.045				1	050					1055		
45							•										•
	acg	tcg	tat	atg	acc	acg	cca	tgc	aag	ctc	att	gat	cta	gga	aat	ggc	3216
				Met													
50				.060				-	065			-		070		-	

	ate	att	ccg	gct	ctc	gag	aac	cgg	gcg	aca	cct	tca	cto	gct	gad	aac	3264
5	Met	Ile	Pro	Ala	Leu	Glu	Asn	Arg	Ala	Thr	Pro	Ser	Leu	Ala	Asp	Asn	
			1075	;				1080	)				1085				
10	acg	aaa	tct	ttc	gaa	att	ctg	ctc	gcc	gaa	gac	aac	acc	gtc	aac	caa	3312
						Ile											
15		1090					1095					1100					
	cga	tta	gça	gtg	aaa	att	ctc	gag	aag	tat	cac	cat	gtg	gta	aca	gtg	3360
20						Ile											
	110	5	-			1110					1115					1120	
25																	
	gtt	ggt	aac	ggc	tgg	gaa.	gct	gtc	aaa	gcc	gtc	caa	agc	aag	aaa	ttc	3408
	Val	Gly	Asn	Gly	Trp	Glu	Ala	Va1	Lys	Ala	Val	Gln	Ser	Lys	Lys	Phe	
30					1125					1130					1135		
												•			•		
 <b>3</b> 5	gat	gtc	att	ctt	atg	gat	gta	caa	atg	ccg	atc	atg	gga	ggc	ttc	gaa	3456
-	Asp	Val	Ile	Leu	Met	Asp	Val	Gln	Met	Pro	Ile	Met	Gly	Gly	Phe	Glu	•
			1	1140				1	1145				1	150			
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45	Ala	Thr	Gly	Lys	Ile	Arg	Glu	Tyr	Glu	Arg	Gly	Ile	Gly	Ser	His	Arg	
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50	aca	ccc	atc	att	gct	cta	acg	gcc	cac	gcc	atg	atg	ggt	gac	cga	gag	3552
						Leu			•								
55													-	•	-		

		1170					1175					1180					
5																	
3	aag	tgt	atc	caa	gct	cag	atg	gac	gag	tat	tte	tcc	aaa	ccc	tte	CSF	3600
																_	0000
10			110	OIN	•			voh	010			Ser	Lys	FFO	Leu	GIN	
	118	5				1190					1195				•	1200	
																	•
15	caa	aac	cat	ctc	atc	cag	acg	atc	ctc	aaa	tgc	gcg	acg	ctc	ggc	ggc	3648
	Gln	Asn	His	Leu	Ile	Gln	Thr	Ile	Leu	Lys	Cys	Ala	Thr	Leu	Gly	Gly	
					1205					1210			•		1215		
20																	
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0.5									•			Ala					
25				1220	2,3	7150	мв			UIU	Leu	NIG			ALA	. GIU	
			•	1220				•	1225				•	1230			
30						•			•								•
	acg	aaa	tcg	aag	cac	aag	gag	ggg	gga	cag	ggt	ctg	cta	cga	ccc	aca	3744
•	Thr	Lys	Ser	Lys	His	Lys	Glu	Gly	Gly	Gln	Gly	Leu	Leu	Arg	Pro	Thr	
35		1	1235				1	240				1	245				•
							•		-								
	ctc	gag	agc	cgc	tca	ttc	aca	agt	cga	gaa	cct	ctg	ttg	gga	aat	ggc	3792
40							-					Leu					
		250					255					260		,		,	
45		200					. 200	•			4	200					
45															•		
								_				gat		_		_	3840
50	Lys	G1u	Ser	Pro	Ala	Ile	Leu	Ala	Thr	Asp	Glu	Asp	Pro	Leu	Ala	Arg	
	1265				1	270				1	275				1	280	

	gca cgt ctt gac ctc tct gat atg cga agt ctt acc aac taa	3882
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20	<213> Artificial Sequence	
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5	<400> 44	
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30	<400> 45 -	
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40	<210> 46	
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	<212> DNA	
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<b>35</b>	•		
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	Ala Gln M	det Asp	Glu Ty	r Leu	Ser	Lys	Pro	Leu	Lys	Xaa	Asn	Gln	Leu
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•							-					•		,			
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25		•											•				
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30			Asn		• •												108
	1111	116	7311	Lys	245	· aı	vəh		Leu	250	Ulu	rue	nia	Set		val .	
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35 -	222		***										_ • -				016
		•	ttg														816
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	ate	gtg	gga	cag	ctc	CFF	acc	ttc	gca	acø	<b>ភ</b> ព ម	øtt	ac a	909	ata	tog	1056
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	щес	, Tal	ĠŢĀ		Leu	VIR	ш	rne.		Inr	GIU	Val	inr			26L	
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				÷													
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•			•		atg												1152
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		:														,	
	gca	aac	aat	ctc	act	gcc	cag	gtg	cga	gat	att	gcg	gag	gtg	aca	aca	1200
50	Ala	Asn	Asn	Leu	Thr	Ala	G1n	Val	Arg	Asp	Ile	Ala	Glu	Val	Thr	Thr	
	385					390			_	•	395					400	
56								•									
55																	

5		gco	gtg	gee	cga	ggc	gac	cto	acg	cag	cag	gtt	aaa	gcg	caa	tgt	aag	1248	
		Ala	Val	Ala	Arg	Gly	Asp	Leu	Thr	Glm	Gln	Val	Lys	Ala	G1n	Cys	Lys		
						405					410					415			
10					•						٠.					÷			
		ggg	gag	atc	ctg	gcc	ttg	aaa	acc	acc	atc	aac	tcc	atg	gtg	cac	cag	1296	
15		Gly	Glu	Ile	Leu	Ala	Leu	Lys	Thr	Thr	Ile	Asn	Ser	Met	Val	His	G1n		
					420		•			425	•				430				
20		cta	cgg	caa	ttc	gcg	cat	gaa	gtc	acc	aag	atc	gcg.	cgt	gag	gtc.	ggg	1344	
									•					Arg					
25		`		435					440		-			. <b>44</b> 5					
			•					-											
		aca	gaa	ggt	CFC	cta	ggt	PFA	caa.	eca.	aca	øtt	cac	gga	atc	as a	aac	1392	
30														Gly			-	1032	•
			450	01,		200	. d.T.)	455		VIT	110	VAI	460	ULY	101	Olu	Gly		
· 35			400	•				730	•				400		,				
	-	363	+~~	000		**-					:							1440	
			•															. 1440	
40			пр	Lys	ASP	Leu		GIU	Asn	Val	Asn		Met	Ala	Met	Asn			
		465					470				• •	475					480		
45					· .													•	
45	•									•				gcc -				1488	
		Thr	Thr	G1n	Val	Arg	G1u	Ile	Ala	Glu.	Val	Thr	Thr	Ala	Val	Ala	Gln		
50						485					490					495			
		•															•		
		gga	gat	ctc	agc	aaa	aag	gtc	gag	gcc	gaa	gtċ	aag	ggt	gaa	att	ttg	1536	
55																			

	Gly	Asp	Leu	Ser	Lys	Lys	Val	Glu	Ala	Glu	Val	Lys	Gly	r Glu	Ile	Leu	
5		٠		500					505	;				510	١		
									•						•		
	gco	ttg	aag	agc	acc	atc	aat	tcc	atg	gtt	gac	cgt	ctg	ggt	acg	ttt	1584
10	Ala	Leu	Lys	Ser	Thr	Ile	Asn	Ser	Met	Val	Asp	Arg	Leu	Gly	Thr	Phe	
			515					520			•		525	i			
15		•															
	gct	ttc	gag	gtt	agc	aaġ	gtc	gcg	aga	gaa	gtc	gga	acc	gaa	gga	gtt	1632
	Ala	Phe	Glu	Val	Ser	Lys	.Val	Ala	Arg	Glu	Va1	G1y	Thr	Glu	Gly	Val	
20		530					535					540					
•							•		•						-	*	
25	ttg	ggc	gga	caa	gca	gag	gtt	gcc	aat	gtc	gaa	gga	aaa	tgg	aaa	gat	1680
	Leu	Gly	Gly	Gln	Ala	G1u	Val	Ala	Asn	Val	Glu	G1y	Lys	Trp	Lys	Asp	
	<b>54</b> 5					<b>550</b>					555	-			•	560	
<b>30</b>				•												•	
	ctt	acc	gac	aat	gtc	aac	acc	atg	gcc	aac	aac	ttg	act	ggt	cag	gtg	1728
35	Leu	Thr	Asp	Asn	Va1	Asn	Thr	Met	Ala	Asn	Așn	Leu	Thr	Gly	Gln	Val	
					565					570					575		
40																	•
40	cgg	agc.	att	tca	gac	gtc	aca	cag	gcc	att	gca	cgc	ggt	gac	atg	agc	1776
	Arg	Ser	Ile	Ser	Asp	Va1	Thr	Gln	Ala	Ile	Ala	Arg	Gly	Asp	Met	Ser	
45				580				•	585					590			
				•		•			•								٠
50	cag	cga	atc	aag	gtg	cac	gct	cag	gga	gag	att	cag	aca	ttg	aag	gac	1824
50	Gln	Arg	Ile	Lys	Val	His	Ala	G1n	Gly	Glu	Ile	Gln	Thr	Leu	Lys	Asp	
			595					600					605				
55																	

5	acg at	c aac	gac atg	gtg	acg	cga	ctg	gac	gct	tgg	tca	ctc	gcg	gtg	1872
	Thr Il	e Asn	Asp Met	Val	Thr	Arg	Leu	Asp	Ala	Trp	Ser	Leu	Ala	Val	
	. 61	0			615					620					
10															•
	aag cg	g gtg	get egt	gac	gtc	ggt	gtc	gac	ggc	aag	atg	ggt	gga	cag	1920
15	_		Ala Arg												•
	625			630		-		•	635	-•			,	640	
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20	700 CO	a est o	505 050					· 			14.				
			gaa ggc												1968
	Ala GI	1 AST	Glu Gly	110	lhr	GTA	Arg		Lys	Glu	Ile	Thr		Asp	
25			645					650					<b>6</b> 55		
30	gtg aad	att	atg gct	caa	aat	ttg	acc	tcg	caa	gtg	aga	gct	ttt	gcc	2016
	Val Ası	Ile	Met Ala	Gln	Asn	Leu	Thr	Ser	Gln	Val	Arg	Ala	Phe	Ala	
			660				665					670			
35					. ,									•	
-	gac att	acc	cac gcg	gcc	atg	aaa	gga	gat	ttc	acc	aag	atg	atc	aat	2064
	Asp Ile	Thr	His Ala	Ala	Met	Lys	Gly	Asp	Phe	Thr	Lys	Met	Ile	Asn	
40 .		675	•			680					685				
45	etc gaa	ge 9 3	tct ggc	gaa .	atø.	990	ga er	cta	920	220	220	ato	336	320	2112
				•		-						•		•	2112
			Ser Gly			ASII	GIU	Leu			Lys	He	ASN .	Lys	•
50	690			•	695		_			700					•
			•												
	atg <sub>,</sub> gtc	ctc	aac ttg	cgc (	gaa	agt .	atc	cag	aag	aac	aat	caa	gca	aga	2160
55												:			

		Met	. Val	Leu	Asu	Leu	Arg	G]u	Ser	I1e	Gln	Lуз	Λen	Asn	Gln	Ala	Arg	
5		705	;				710	)				715		٠			720	
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		gag	gco	gcc	gag	ttg	gcc	aac	aag	acg	asa	tcg	gag	ttc	ctg	gca	aac	2208
10		Glu	Ala	Ala	G1u	Leu	Ala	Asn	Lys	Thr	Lys	Ser	Glu	Phe	Leu	Ala	Asn	
						725					730					735		
15	•						-	•										
		atg	tcc	cac	gag	att	cga	aca	cct	atg	aac	gga	atc	atc	gga	atg	aca	2256
	•							Thr						•				
20					740					745					750		,	
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25		cag	ctt	acc	ttg	gac	acc	gag	ctt	gag	cag	aac	caa	cgg	gac	atg	ctc	2304
								G1u				-						
				755					760					765				
30					-					٠								
		aac	atc	.gtc	ttc	tcg	ctc	gcc	aac	agc	tta	ctg	acg	att	att	gat	gac	2352
 35								Ala		•								
	-		770					775	•		•		780					
			٠															
40		atc	ttg	gac	att	tcc	aag	att	gaa	gca	aat	cgc	atg	atc	cta	gag	gaa	2400
		Ile	Leu	Asp	Ile	Ser	Lys	Ile	Glu	Ala	Asn	Arg	Met	Ile	Leu	Glu	G1u	
45		785					790		•		•	795					800	
				•					•									`
		gag	ccg	ttc	tca	ctg	cga	ggt	cţc	gtc	ttc	aac	agc	tta	aag	tca	ctt	2448
50								Gly										
						805					810					815		
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5	gca	gto	cga	gco	aac	gag	aag	gac	ato	ago	tte	gtg	tat	gat	acc	gac	2496
	Ala	a Val	Arg	, Ala	Asn	Glu	Lys	Asp	Ile	e Ser	Leu	ı Val	Tyr	Asp	Thr	. Asp	
				820	)				825	,				830	)		•
10																	
	ago	tca:	gte	ccc	gac	tac	atc	gtg	FFC	. gar	tro	tto	ras	ctt	649	cag	2544
•															_	_	2077
15	DET	Det			, veh	IÀT	TIE			ASP	Ser	rne			Arg	Gln	
			835	•				840					845				
20					•												
20	ato	att	ctc	aat	ctc	gcc	ggc	aac	gcc	atc	aaa	ttc	acc	gag	cac	ggg	2592
	Ile	Ile	Leu	Asn	Leu	Ala	Gly	Asn	Ala	Ile	Lys	Phe	Thr	G1u	His	Gly	
25		850					855					860					
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•	gaa	gtg	cøt	gtt	aag	ata	ttc	tet	<b>72</b> 0	Cac	act		C (7.2	+ 00	200	ga+	2640
30					•											•	2040
			ив	141	Lys		rne	Ser	ASD	nis		1111	Arg .	Cys	ınr		
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35						•								•			
	agt	gag	gtt	gtc	gtc	aaa	ttc	gcc	gtc	tcc	gat	act	ggt	att	ggc	atc	2688
	Ser	G1u	Val	Val	Val	Lys	Phe	Ala	Va1	Ser	Asp	Thr	Gly	Ile	Gly	Ile	
<sub>.</sub> 40					885					890					895		
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45	cac	tee	aac	220	ttg	ast.	tta	atr	ttc	<b>42</b> C	2011		025	000	aat		2736
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	nis	ser	ASR		Leu	Asp	Leu			Asp	inr	Phe	GĻn	Gln	Ala	Asp	
50				900		٠			905					910	•		·
			•						•								
	EEE	tċg	acc	aca	cgg	aag	ttc	gga	ggt	act	gga	ttg	ggc	ctg	tcg	atc	2784
55																	

	Gly	/ Ser	The	Thi	Arg	Lys	Phe	Gly	G13	Thr	Gly	Let	Gly	/ Let	ı Ser	· Ile	
5			915	;				920	)				925	5			
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10	tct	cgg	aga	cte	gtg	act	tte	atg	cgt	ggc	aag	atg	tgg	gto	gaa	tca	2832
	Sex	Arg	Arg	Leu	Val	Thr	Leu	Met	Arg	Gly	Lys	Met	Trp	Val	Glu	Set	
	•	930	)		•		935	i				940					
15							-										
	aat	tat	ggc	tca	ggc	agc	aca	ttc	ttc	ttc	acc	tgk	gtt	gta	cgg	ctg	2880
20	Asn	Tyr	Gly	Ser	Gly	Ser	Thr	Phe	Phe	Phe	Thr	Xaa	Val	Val	Arg	Leu	
	945					950					955			٠	٠.	960	•
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25	ggc	aat	ccg	gat	gtt	gca	aaa	atc	atg	cca	caa	cta	cag	cag	tat	cga	2928
	Gly	Asn	Pro	Asp	Va1	Ala	Lys	Ile	Met	Pro	Gln	Leu	Gln	Gln	Тут	Arg	•
30					965			•	•	970					975		
-																	
	aag	cac	aac	gtg	ctc	ttt	gtc	gaç	aac	ggt	aat	acg	gac	agt	tcg	gag	2976
<b>35</b> –	Lys	His	Asn	Val	Leu	Phe	Va1	Asp	Asn	Gly	Asn	Thr	Asp	Ser	Ser	G1u	•
				980				٠	985					990			
40																	
	gag	atc	gcg	gct	ggc	atc	cga	gct	ttg	gat	ctg	gtc	cct	tgt	gtg	gtg	3024
	Glu	Ile	Ala	Ala	G1y	Ile	Arg	Ala	Leu	Asp	Leu	Val	Pro	Cys	Val	Val	
45			995				1	.000			-	1	.005				•
50						•											
	ggc	aag	gga	aag	gtt	cct	cac	tcc	gaa	atc	agc	cca	gac	gac	cag	tac	3072
55	Gly	Lys	Gly	Lys	Val	Pro	His	Ser	Glu	Ile	Ser	Pro	Asp	Asp	Gln	Tyr	•
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	1010		1015	1020	l	
5						
	gac tgc	gtg atc atc	gat aac agc	gag acg gct cag	aag ttg cgc agc	3120
10	Asp Cys	Val Ile Ile	Asp Asn Ser	Glu Thr Ala Gln	Lys Leu Arg Ser	•
10	1025	1	1030	1035	1040	
15	ttg gaa	cgt ttc aag	tac att ccc	atc gtc atg gtg	gcg ccg gcc atc	3168
				Ile Val Met Val		
		1045		1050	1055	
20			· .		2000	
	tcg gtg	aac ttc aag	acc gcg ttg	gag aac gga atc	tca agc tac atg	3216
25	Ser Val	Asn Phe Lys	Thr Ala Leu	Glu Asn Gly Ile	Ser Ser Tyr Met	
	•	1060	1	1065	1070	
	•					
30	act acg	cca tgc ctt	cca atc gac	ctg ggc aat gct	ctg gtg ccc gca	3264
	Thr Thr	Pro Cys Leu	Pro Ile. Asp	Leu Gly Asn Ala	Leu Val Pro Ala	
35	1	075	1080	٠.	1085	
		•	·			
40	ctc gag	ggc cgc gca	gca ccc atg	tca gcc gac cac	agt cgg aca ttc	3312
	Leu Glu (	Gly Arg Ala	Ala Pro Met	Ser Ala Asp His	Ser Arg Thr Phe	
	1090		1095	1100		
45		•	•			
	gat atc	ctc ctc gca	gaa gac aac	gcg gtg aat caa	aaa ctc gcc gtc	3360
50	Asp Ile I	Leu Leu Ala	Glu Asp Asn	Ala Val Asn Gln	Lys Leu Ala Val	
50	1105	1.	110	1115	1120	
			•			

Lys Ile Leu Thr Lys His Asn His Thr Val Thr Val Ala Asn Asn Gly  1125 1130 1135  10  ctt gaa gcc ttt gaa gcg att cgc aag aag cgc ttc gat gtc gtt ctc Leu Glu Ala Phe Glu Ala Ile Arg Lys Lys Arg Phe Asp Val Val Leu  1140 1145 1150  atg gac gtg caa atg ccc gtc atg gga ggg ttc gaa gcg acg gcc aag  20  Met Asp Val Gln Met Pro Val Met Gly Gly Phe Glu Ala Thr Ala Lys  1165 1160  1165  25  att cgc gaa tac gaa cgc act cac gag cta gca cgt tcg ccc att atc 1170 1175 1180  36  gcc ctc acc gca cac gcc atg ctt ggc gac cgc gag aag tgt atc caa Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln 1185 1190 1195 1200  gcg caa atg gac gag tat ctc tcc aaa ccc ctc aag ycc aat cag ctc 3648  Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210  1215  att cag acg atc ctg aaa tgt gog acc cta ggc ggt gcg tta ctt gac 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp		aag	atc	ctg	acc	aag	cạc	aac	cac	aca	gtg	aca	gtc	gcc	aac	aac	ggc	3408
ctt gaa gcc ttt gaa gog att cgc aag aag cgc ttc gat gtc gtt ctc  Leu Glu Ala Phe Glu Ala Ile Arg Lys Lys Arg Phe Asp Val Val Leu  1140 1145 1150  atg gac gtg caa atg ccc gtc atg gga ggg ttc gaa gcg acg gcc aag 3504  Met Asp Val Gln Met Pro Val Met Gly Gly Phe Glu Ala Thr Ala Lys  1166 1160 1165  att cgc gaa tac gaa cga act cac gag cta gca cgt tcg ccc att atc 3552  Ile Arg Glu Tyr Glu Arg Thr His Glu Leu Ala Arg Ser Pro Ile Ile  1170 1175 1180  36 gcc ctc acc gca cac gcc atg ctt ggc gac cgc gag aag tgt atc caa 3600  Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln  1185 1190 1195 1200  gcg caa atg gac gag tat ctc tcc aaa ccc ctc aag ycc aat cag ctc 3648  Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gin Leu  1205 1210 1215  att cag acg atc ctg aaa tgt gcg acc cta ggc gst gcg tta ctt gac 3696  Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp	5	Lys	Ile	Leu	Thr	Lys	His	Asn	His	Thr	Val	Thr	Val	Ala	Asn	Asn	Gly	
Leu Glu Ala Phe Glu Ala Ile Arg Lys Lys Arg Phe Asp Val Val Leu  1140 1145 1150  atg gac gtg caa atg ccc gtc atg gga ggg ttc gaa gcg acg gcc aag 3504  Met Asp Val Gln Met Pro Val Met Gly Gly Phe Glu Ala Thr Ala Lys 1155 1160 1165  25  att cgc gaa tac gaa cga act cac gag cta gca cgt tcg ccc att atc 3552  Ile Arg Glu Tyr Glu Arg Thr His Glu Leu Ala Arg Ser Pro Ile Ile 1170 1175 1180  36  gcc ctc acc gca cac gcc atg ctt ggc gac cgc gag aag tgt atc caa 3600  Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln 1185 1190 1195 1200  46  gcg caa atg gac gag tat ctc tcc aaa ccc ctc aag ycc aat cag ctc 3648  45  Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210 1215  50  att cag acg atc ctg aaa tgt ggg acc cta ggc ggt gg tta ctt gac 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp						1125					1130					1135		
Leu Glu Ala Phe Glu Ala Ile Arg Lys Lys Arg Phe Asp Val Val Leu  1140 1145 1150  atg gac gtg caa atg ccc gtc atg gga ggg ttc gaa gcg acg gcc aag 3504  Met Asp Val Gln Met Pro Val Met Gly Gly Phe Glu Ala Thr Ala Lys 1155 1160 1165  25  att cgc gaa tac gaa cga act cac gag cta gca cgt tcg ccc att atc 3552  Ile Arg Glu Tyr Glu Arg Thr His Glu Leu Ala Arg Ser Pro Ile Ile 1170 1175 1180  36  gcc ctc acc gca cac gcc atg ctt ggc gac cgc gag aag tgt atc caa 3600  Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln 1185 1190 1195 1200  46  gcg caa atg gac gag tat ctc tcc aaa ccc ctc aag ycc aat cag ctc 3648  45  Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210 1215  50  att cag acg atc ctg aaa tgt ggg acc cta ggc ggt gg tta ctt gac 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp																		
atg gac gtg caa atg ccc gtc atg gga ggg ttc gaa gcg acg gcc aag 3504  Met Asp Val Gln Met Pro Val Met Gly Gly Phe Glu Ala Thr Ala Lys 1155 1160 1165  att cgc gaa tac gaa cga act cac gag cta gca cgt tcg ccc att atc 3552  Ile Arg Glu Tyr Glu Arg Thr His Glu Leu Ala Arg Ser Pro Ile Ile 1170 1175 1180  36 gcc ctc acc gca cac gcc atg ctt ggc gac ogc gag aag tgt atc caa 3600 Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln 1185 1190 1195 1200  gcg caa atg gac gag tat ctc tcc aaa ccc ctc aag ycc aat cag ctc 3648  Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210 1215  att cag acg atc ctg aaa tgt gcg acc cta ggc ggt gcg tta ctt gac 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp	10	ctt	gaa	gcc	ttt	gaa	gcg	att	cgc	aag	aag	cgc	ttc	gat	gtc	gtt	ctc	3456
atg gac gtg caa atg ccc gtc atg gga ggg ttc gaa gcg acg gcc aag 3504  Met Asp Val Gln Met Pro Val Met Gly Gly Phe Glu Ala Thr Ala Lys  1166  1160  1165  25  att cgc gaa tac gaa cga act cac gag cta gca cgt tcg ccc att atc 3552  Ile Arg Glu Tyr Glu Arg Thr His Glu Leu Ala Arg Ser Pro Ile Ile 1170  1175  1180  36  gcc ctc acc gca cac gcc atg ctt ggc gac cgc gag aag tgt atc caa Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln 1185  1190  1195  1200  gcg caa atg gac gag tat ctc tcc aaa ccc ctc aag ycc aat cag ctc 3648  Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205  1210  1215  att cag acg atc ctg aaa tgt gcg acc cta ggc ggt gcg tta ctt gac 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp		Leu	Glu	Ala	Phe	Glu	Ala	Ile	Arg	Lys	Lys	Arg	Phe	Asp	Val	Val	Leu	
Met Asp Val Gln Met Pro Val Met Gly Gly Phe Glu Ala Thr Ala Lys 1166 1160 1165  att cgc gaa tac gaa cga act cac gag cta gca cgt tcg ccc att atc 3552  Ile Arg Glu Tyr Glu Arg Thr His Glu Leu Ala Arg Ser Pro Ile Ile 1170 1175 1180  3600  Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln 1185 1190 1195 1200  46  gcg caa atg gac gag tat ctc tcc aaa ccc ctc aag ycc aat cag ctc 3648  45  Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210 1215  att cag acg atc ctg aaa tgt gcg acc cta ggc ggt gcg tta ctt gac 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp	15				1140	•			:	1145					1150			
Met Asp Val Gln Met Pro Val Met Gly Gly Phe Glu Ala Thr Ala Lys 1166 1160 1165  att cgc gaa tac gaa cga act cac gag cta gca cgt tcg ccc att atc 3552  Ile Arg Glu Tyr Glu Arg Thr His Glu Leu Ala Arg Ser Pro Ile Ile 1170 1175 1180  3600  Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln 1185 1190 1195 1200  46  gcg caa atg gac gag tat ctc tcc aaa ccc ctc aag ycc aat cag ctc 3648  45  Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210 1215  att cag acg atc ctg aaa tgt gcg acc cta ggc ggt gcg tta ctt gac 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp		•										•		•				
att cgc gaa tac gaa cga act cac gag cta gca cgt tcg ccc att atc 3552  Ile Arg Glu Tyr Glu Arg Thr His Glu Leu Ala Arg Ser Pro Ile Ile  1170 1175 1180  gcc ctc acc gca cac gcc atg ctt ggc gac gg gag aag tgt atc caa 3600  Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln  1185 1190 1195 1200  gcg caa atg gac gag tat ctc tcc aaa ccc ctc aag ycc aat cag ctc 3648  Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu  1205 1210 1215  att cag acg atc ctg aaa tgt gcg acc cta ggc ggt gcg tta ctt gac 3696  Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp	20	atg	gaç	gtg	caa	atg	ccc	gtc	atg	gga	ggg	ttc	gaa	gcg	acg	gcc	aag	3504
att cgc gaa tac gaa cga act cac gag cta gca cgt tcg ccc att atc 3552  Ile Arg Glu Tyr Glu Arg Thr His Glu Leu Ala Arg Ser Pro Ile Ile  1170 1175 1180  gcc ctc acc gca cac gcc atg ctt ggc gac cgc gag aag tgt atc caa 3600  Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln  1185 1190 1195 1200  gcg caa atg gac gag tat ctc tcc aaa ccc ctc aag ycc aat cag ctc 3648  Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu  1205 1210 1215  att cag acg atc ctg aaa tgt gcg acc cta ggc ggt gcg tta ctt gac 3696  Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp	20	Met	Asp	Val	Gln	Met	Pro	Val	Met	Gly	Gly	Phe	Glu	Ala	Thr	Ala	Lys	
att cgc gaa tac gaa cga act cac gag cta gca cgt tcg ccc att atc  Ile Arg Glu Tyr Glu Arg Thr His Glu Leu Ala Arg Ser Pro Ile Ile  1170 1175 1180  gcc ctc acc gca cac gcc atg ctt ggc gac cgc gag aag tgt atc caa 3600 Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln 1185 1190 1195 1200  gcg caa atg gac gag tat ctc tcc aaa ccc ctc aag ycc aat cag ctc Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210 1215  att cag acg atc ctg aaa tgt gcg acc cta ggc ggt gcg tta ctt gac 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp			:	1155					1160				•	1165				•
Ile Arg Glu Tyr Glu Arg Thr His Glu Leu Ala Arg Ser Pro Ile Ile  1170 1175 1180  35 gcc ctc acc gca cac gcc atg ctt ggc gac cgc gag aag tgt atc caa 3600 Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln 1185 1190 1195 1200  40 gcg caa atg gac gag tat etc tcc aaa ccc ctc aag ycc aat cag etc 3648  45 Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210 1215  50 att cag acg atc ctg aaa tgt gcg acc cta ggc ggt gcg tta ctt gac 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp	25												٠					
gcc ctc acc gca cac gcc atg ctt ggc gac cgc gag aag tgt atc caa 3600 Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln 1185 1190 1195 1200  gcg caa atg gac gag tat etc tcc aaa eec etc aag ycc aat cag etc 3648 Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210 1215  att cag acg atc etg aaa tgt gcg acc eta ggc ggt gcg tta ett gac 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp			•			_	-			•		-						3552
gcc ctc acc gca cac gcc atg ctt ggc gac cgc gag aag tgt atc caa 3600 Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln 1185 1190 1195 1200  gcg caa atg gac gag tat ctc tcc aaa ccc ctc aag ycc aat cag ctc 3648 Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210 1215  att cag acg atc ctg aaa tgt gcg acc cta ggc ggt gcg tta ctt gac 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp	30			Glu	Tyr	Glu			His	Glu	Leu			Ser	Pro	Ile	Ile	
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Ala Leu Thr Ala His Ala Met Leu Gly Asp Arg Glu Lys Cys Ile Gln 1185 1190 1195 1200  geg caa atg gae gag tat etc tec aaa eec etc aag yee aat eag etc 3648  Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210 1215  att eag aeg ate etg aaa tgt geg aec eta gge ggt geg tta ett gae 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp	_	•					•									•		2222
gcg caa atg gac gag tat etc tec aaa eec etc aag yee aat eag etc 3648  Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210 1215  att eag acg atc etg aaa tgt geg ace eta gge ggt geg tta ett gac 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp	<b>35</b>																	3600
geg caa atg gae gag tat etc tec aaa eec etc aag yee aat eag etc 3648  Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210 1215  att eag aeg ate etg aaa tgt geg aec eta gge ggt geg tta ett gae 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp				Ihr	Ala			Met	Leu	GIA	_	_	GIU	Lys	Cys			
Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210 1215  att cag acg atc ctg aaa tgt gcg acc cta ggc ggt gcg tta ctt gac 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp	40	110	•				1190				1	1195					1200	
Ala Gln Met Asp Glu Tyr Leu Ser Lys Pro Leu Lys Xaa Asn Gln Leu 1205 1210 1215  att cag acg atc ctg aaa tgt gcg acc cta ggc ggt gcg tta ctt gac 3696 Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp				n+ ~	<b>~~</b>		***	a±a	+	222		ata	222		+	005	***	3640
1205 1210 1215  50  att cag acg atc ctg aaa tgt gcg acc cta ggc ggt gcg tta ctt gac 3696  Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp	45				•													. 3040
att cag acg atc ctg aaa tgt gcg acc cta ggc ggt gcg tta ctt gac 3696  Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp	45	nia	0111	Met	_		LYL	Leu	Jer			Leu	Lys	Ada -			Leu	•
att cag acg atc ctg aaa tgt gcg acc cta ggc ggt gcg tta ctt gac 3696  Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp						200				4	.210				•	1213		
Ile Gln Thr Ile Leu Lys Cys Ala Thr Leu Gly Gly Ala Leu Leu Asp	50	att	Cag	acø	atc	cte	222	tøt	g c g	acc	cta	ggc	oot	aca	tta	ctt	gac.	3696
		•																3300
55	55			_			-,-	-,-				• •						

			:	1220	1225					5 1230							
5																	
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	Arg	Arg	Asn	Asp	Gly	Arg	G1y	Leu	Leu	Met	Glu	Glu	Asp	Lys	Pro	Yal	
10		,1	1235					1240				C	1245	•			
					٠		•										
15	tct	gat	aat	tcg	agt	ctt	cct	gca	gat	cac	aat	cgg	ttg	ctc	acg	ccc	3792
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	Gly	Leu	Glu	Ser	Pro	Ala	Ile	Val.	Thr	Asp	Asp	Gln	Asp	Asp	Pro	Met	
- 35				. 1	<b>28</b> 5	. •			1	1290					295		
-		•															
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	Thr	Asp	Gly	Asp	Phe	Thr	Arg	Phe	Ile	Thr	Val	Glu	Ala	Ser	Gly	Glu
50					885					890		٠			895	
	Met	Asp	Ser		Lys	Thr	Gln	Ile		Gln	Met	Val	Tyr		Leu	Arg
55				900					905					910		

	Glu	Ser	Ile	Gln	Arg	Asn	Thr	Ala	Ala	Arg	Glu	Ala	Ala	Glu	Leu	Ala
5			915					920					925		•	
	Asn	Arg	Ser	Lys	Ser	G1u	Phe	Leu	Ala	Asn	Met	Ser	His	Glu	Ile	Arg
10		930					935					940				
	Thr	Pro	Met	Asn	Gly	Ile	Ile	Gly	Met	Thr	Asp	Leu	Thr	Leu	Asp	Thr
	945					950					955					960
15	Glu	Leu	Thr	Arg	Thr	Gln	Lys	Glu	Asn	Leu	Leu	Leu	Val	His	Gln	Leu
					965					970					975	
20	Ala	Lys	Ser	Leu	Leu	Leu	·Ile	Ile	Asp	Asp	Ile	Leu	Asp	Ile	Ser	Lys
				980					985					990		
	Ile	Glu	Ala	Gly	Arg	Met	Thr	Met	Glu	Gln	Val	Thr	Tyr	Ser	Leu	Arg
25			995			-	;	1000		٠		. 1	1005			
	Gly	Thr	Àla	Phe	Gly	Ile	Leu	Lys	Thr	Leu	Val	Val	Arg	Ala	His	Gln
30	:	1010			٠	i.	1015				;	1020	•			
. <del>-</del>	Gln	Asn	Leu	Asn	Leu	Phe	Tyr	Glu	Val	Asp	Pro	G1u	Ile	Pro	Asp	Gln
	1025	5			]	030					035				1	040
<b>35</b> -	Val	Ile	Gly	Asp	Ser	Leu	Arg	Leu	Arg	Gln	Val	Ile	Thr	Asn	Leu	Val
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40	Gly	Asn	Ala	Ile	Lys	Phe	Thr	Pro	Ser	Lys	Pro	Asn	Lys	Lys	Gly	Met
			1	060			•	1	065			•	1	070		
45	Val	Cys	Leu	Ser	Cys	Lys	Leu	Ile	Ser	Met	Asp	Glu	Gin	Asn	Val	Thr
45		1	.075				-1	080				1	085			
	Val	Arg	Phe	Cys	Va1	Glu	Asp	Thr	Gly	Ile	Gly	Ile	Lys	Gln	Asp	Lys
50	1	090	•			1	095				1	100				
	Leu	Ala	Ile	Ile	Phe	Asp	Thr	Phe	Cys	Gln	Ala	Asp	Gly	Ser	Thr	Thr
55	1105				1	110				1	115				1	120

£	Arg Glu	Tyr	Gly Gl	y Thr	· Gly	Leu	G1y	Leu	Ser	Ile	Ser	Lys	Arg	Leu
5			112	5				1130					1135	
	Val Ser	Leu !	Met As	n Gly	Gln	Met	Trp	Val	Glu	Ser	Glu	Val	Gly	Val
10		1	140				1145					1150		
	Gly Ser	Arg	Phe Ty	r Phe	Thr	Ile	Thr	Ala	Glu	Ile	Ser	Arg	Pro	Asn
15		1155		÷		1160			٠	•	1165			
,,,	Met Ala	Gln :	Ser Le	u Gln	Lys	Val	Ala	Ile	Tyr	Lys	Glu	Arg	Thr	Ile
	1170				1175					1180				
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	1185			1190					1195					1200
25	Glu Glu	Leu	Gln Le	u Arg	Pro	Phe	Val	Val	Arg	Asp	Ile	Ser	G1n	Val
			120	5				1210					1215	
	Ala Asp	Lys I	Ala Ly	s Ile	Pro	Phe	Ile	Asp	Thr	Val	Ile	Val	Asp	Ser
30		12	220			]	1225				:	1230		
30	Leu Glu			ı Lys	Leu			Leu	Asp	His			Tyr	Thr
30 				ı Üys				Leu	Asp				Tyr	Thr
-/		Val 1 1235	Thr Gl			Arg 1240	Glu			]	Leu 1245	Arg		
-/		Val 1 1235	Thr Gl	ı Thr		Arg 1240	Glu		Arg	]	Leu 1245	Arg		
-/	Pro Ala	Val 1 1235 Val 1	Thr Gl	ı Thr	Pro 1255	Arg 1240 Val	Glu Met	Pro	Arg	Leu (260	Leu 1245 Asn	Arg Leu	Thr	Trp
35 35	Pro Ala 1250	Val 1 1235 Val 1	Thr Gl	ı Thr	Pro 1255	Arg 1240 Val	Glu Met	Pro Val	Arg	Leu (260	Leu 1245 Asn	Arg Leu	Thr	Trp
35 35	Pro Ala 1250 Cys Leu	Val 1 1235 Val I Glu A	Thr Gl Leu Le Asn Ph	1 Thr 2 Ile 1270	Pro 1255 Ser	Arg 1240 Val Gly	Glu Met His	Pro Val	Arg Ala 1275	Leu (260 Thr	Leu 1245 Asn Pro	Arg Leu Ser	Thr Ser	Trp Leu 280
35	Pro Ala 1250 Cys Leu 1265	Val 1 1235 Val I Glu A	Thr Gl Leu Le Asn Ph	Thr Ile 1270	Pro 1255 Ser	Arg 1240 Val Gly	Glu Met His	Pro Val	Arg Ala 1275	Leu (260 Thr	Leu 1245 Asn Pro	Arg Leu Ser Asn	Thr Ser	Trp Leu 280
35	Pro Ala 1250 Cys Leu 1265	Val 1 1235 Val I Glu A	Thr Gl Leu Le Asn Ph Ala Gl	I Thr	Pro 1255 Ser Leu	Arg 1240 Val Gly Ala	Glu Met His Lys	Pro Val Gly	Arg Ala 1275 Leu	Leu 1260 Thr Glu	Leu 1245 Asn Pro	Arg Leu Ser Asn	Thr Ser Ala 295	Trp Leu 280 Ser
35	Pro Ala 1250 Cys Leu 1265 Asp Asp	Val 1 1235 Val I Glu A Glu A	Thr Gl Leu Le Asn Ph Ala Gl	I Thr	Pro 1255 Ser Leu	Arg 1240 Val Gly Ala	Glu Met His Lys	Pro Val Gly	Arg Ala 1275 Leu	Leu 1260 Thr Glu	Leu 1245 Asn Pro Ala	Arg Leu Ser Asn	Thr Ser Ala 295	Trp Leu 280 Ser
35 - 40 45	Pro Ala 1250 Cys Leu 1265 Asp Asp	Val 1 1235 Val I Glu A Glu A	Thr Gl Leu Le Asn Ph Ala Gl 1289 Val Th	I Thr	Pro 1255 Ser Leu Ser	Arg 1240 Val Gly Ala Asp	Glu Met His Lys Val	Pro Val Gly 1290 Ala	Arg Ala 1275 Leu Tyr	Leu (260 Thr Glu	Leu 1245 Asn Pro Ala	Arg Leu Ser Asn Leu 310	Thr Ser Ala 295 Leu	Trp Leu 280 Ser Ala

Phe Gly His Thr Val Gln Ile Ala Glu Asn Gly Gln Phe Ala Val Asp Ala Val Lys Ala Arg Tyr Glu Gln Glu Lys Met Phe Asp Val Ile Leu Met Asp Val Ser Met Pro Phe Met Gly Gly Met Glu Ala Thr Glu Ile Ile Arg Ala Phe Glu Lys Glu Lys Gly Ile Arg Arg Thr Pro Ile Ile Ala Leu Thr Ala His Ala Met Ile Gly Asp Arg Glu Arg Cys Ile Gln Ala Gly Met Asp Glu His Val Thr Lys Pro Leu Arg Arg Thr Asp Leu Val Ser Ala Ile Lys Arg Leu Val Thr Pro His Gly Ala His <210> 69 <211> 4317 <212> DNA (213) Thanatephorus cucumeris <220> <221> CDS <222> (1).. (4317) <400> 69 

Met Ala Gly Thr Thr Gly Gly His Pro Phe Thr Ala His Leu Val Ala  1 5 10 15  gtg ctg agt atc tat gag tta gga ccg gga cga cca gtg cgc gca ctg Val Leu Ser Ile Tyr Glu Leu Gly Pro Gly Arg Pro Val Arg Ala Leu 20 25 30  ccg acc cgg agc tca cat tcc cat tcc tct tcc ggt tcc cgc cat gcg Pro Thr Arg Ser Ser His Ser His Ser Ser Ser Gly Ser Arg His Ala 35 40 45  cgt gcg ctg tct gtg ccg ccg ttc cca cca ccg cca ccg atg tct ccg Arg Ala Leu Ser Val Pro Pro Phe Pro Pro Pro Pro Pro Met Ser Pro 50 55 60	96 144 192
gtg ctg agt atc tat gag tta gga ccg gga cga cca gtg cgc gca ctg  Val Leu Ser Ile Tyr Glu Leu Gly Pro Gly Arg Pro Val Arg Ala Leu  20 25 30  ccg acc cgg agc tca cat tcc cat tcc tct tcc ggt tcc cgc cat gcg Pro Thr Arg Ser Ser His Ser His Ser Ser Ser Gly Ser Arg His Ala  35 40 45  cgt gcg ctg tct gtg ccg ccg ttc cca cca ccg cca ccg atg tct ccg Arg Ala Leu Ser Val Pro Pro Pro Pro Pro Pro Pro Met Ser Pro  50 55 60	144
gtg ctg agt atc tat gag tta gga ccg gga cga cca gtg cgc gca ctg  Val Leu Ser Ile Tyr Glu Leu Gly Pro Gly Arg Pro Val Arg Ala Leu  20 25 30  ccg acc cgg agc tca cat tcc cat tcc tct tcc ggt tcc cgc cat gcg Pro Thr Arg Ser Ser His Ser His Ser Ser Ser Gly Ser Arg His Ala  35 40 45  cgt gcg ctg tct gtg ccg ccg ttc cca cca ccg cca ccg atg tct ccg Arg Ala Leu Ser Val Pro Pro Pro Pro Pro Pro Pro Met Ser Pro  50 55 60	144
Val Leu Ser Ile Tyr Glu Leu Gly Pro Gly Arg Pro Val Arg Ala Leu  20 25 30  ccg acc cgg agc tca cat tcc cat tcc tct tcc ggt tcc cgc cat gcg Pro Thr Arg Ser Ser His Ser His Ser Ser Ser Gly Ser Arg His Ala  35 40 45  cgt gcg ctg tct gtg ccg ccg ttc cca cca ccg cca ccg atg tct ccg Arg Ala Leu Ser Val Pro Pro Phe Pro Pro Pro Pro Pro Met Ser Pro  50 55 60	144
Val Leu Ser Ile Tyr Glu Leu Gly Pro Gly Arg Pro Val Arg Ala Leu  20 25 30  ccg acc cgg agc tca cat tcc cat tcc tct tcc ggt tcc cgc cat gcg Pro Thr Arg Ser Ser His Ser His Ser Ser Ser Gly Ser Arg His Ala  35 40 45  cgt gcg ctg tct gtg ccg ccg ttc cca cca ccg cca ccg atg tct ccg Arg Ala Leu Ser Val Pro Pro Phe Pro Pro Pro Pro Pro Met Ser Pro  50 55 60	144
20 25 30  20 ccg acc cgg agc tca cat tcc cat tcc tct tcc ggt tcc cgc cat gcg Pro Thr Arg Ser Ser His Ser His Ser Ser Ser Gly Ser Arg His Ala 35 40 45  cgt gcg ctg tct gtg ccg ccg ttc cca cca ccg cca ccg atg tct ccg Arg Ala Leu Ser Val Pro Pro Phe Pro Pro Pro Pro Met Ser Pro 50 55 60	
ccg acc cgg agc tca cat tcc cat tcc tct tcc ggt tcc cgc cat gcg Pro Thr Arg Ser Ser His Ser His Ser Ser Ser Gly Ser Arg His Ala  35 40 45  cgt gcg ctg tct gtg ccg ccg ttc cca cca ccg cca ccg atg tct ccg Arg Ala Leu Ser Val Pro Pro Pro Pro Pro Pro Pro Pro Met Ser Pro  50 55 60	
Pro Thr Arg Ser Ser His Ser His Ser Ser Ser Gly Ser Arg His Ala  35  40  45  cgt gcg ctg tct gtg ccg ccg ttc cca cca ccg cca ccg atg tct ccg  Arg Ala Leu Ser Val Pro Pro Phe Pro Pro Pro Pro Pro Met Ser Pro  50  55  60	
25  cgt gcg ctg tct gtg ccg ccg ttc cca cca ccg cca ccg atg tct ccg  Arg Ala Leu Ser Val Pro Pro Pro Pro Pro Pro Pro Met Ser Pro  50  55  60	192
25  cgt gcg ctg tct gtg ccg ccg ttc cca cca ccg cca ccg atg tct ccg  Arg Ala Leu Ser Val Pro Pro Pro Pro Pro Pro Pro Met Ser Pro  50  55  60	192
cgt gcg ctg tct gtg ccg ccg ttc cca cca ccg cca ccg atg tct ccg  Arg Ala Leu Ser Val Pro Pro Phe Pro Pro Pro Pro Pro Met Ser Pro  50 55 60	192
Arg Ala Leu Ser Val Pro Pro Pro Pro Pro Pro Pro Met Ser Pro 50 55 60	192
50 55 60	
ceg aac gca ceg ate gae tac gta gge get get ceg etg eec ega tac	240
Pro Asn Ala Pro Ile Asp Tyr Val Gly Ala Ala Pro Leu Pro Arg Tyr	
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gat gga ccg cgt gac tgg cag acg gat gcg gtc gag cga gca ctg ggc	288
Asp Gly Pro Arg Asp Trp Gln Thr Asp Ala Val Glu Arg Ala Leu Gly	
<b>85</b> 90 95	
50	
cgt gtt gee geg egg atg tac geg gee gag gee eag etg eag gae etg	200
Arg Val Ala Ala Arg Met Tyr Ala Ala Glu Ala Gln Leu Gln Asp Leu	336

				100					105					110				
5																•		
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10	Leu	Ser	Arg	Glu	Ser	Ser	Thr	Ser	Thr	Pro	Asp	Pro	Ala	Leu	Ser	Pro		
		٠	115					120					125					
																•		
15	cgc	tcc	aac	ggc	ctc	aaa	aaa	cgc	aga	gag	aac	ccg	gga	aca	ccc	gat	432	
	Arg	Ser	Asn	Gly	Leu	Lys	Lys	Arg	Arg	G1u	Asn	Pro	Gly	Thr	Pro	Asp		
20		130					135					140						
	gag	cgc	gat	ccg	tgg	cag	act	gtg	cgc	ttt	caa	gag	gtc	ggt	gac	cag	480	
25	Glu	Arg	Asp	Pro	Trp	Gln	Thr	Val	Arg	Phe	G1n	Glu	Va1	Gly	Asp	Gln		
	145					150					155					160		
20					-													
30	gac	atg	oat	ccc	g a ô	cca	<b>79</b> 7	900	cct	at t	<b>acc</b>	ĊĠĊ	C C C	220	#2^	926	528	
																	020	
35	rsh	Met	ASD.	LIO		rro	ASP	Inr	FIO		MIS	WLE	FFO	Lys		Lys		
					165					170		•			175			
		•																
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	Val	Lys	Pro	Gly	Thr	Ile	Asp	Leu	Ser	Thr	Leu	Ser	Gln	Pro	Thr	Pro		
	•			180					185					190				
45						-					•							
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50	Leu	Ser	Lys	Val	Ala	Thr	Asp	Asp	Pro	Val	Leu	Pro	Lys	Pro	Gly	Pro		
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5	Arg	Ser	Ala	Pro	Thr	Ser	Ser	Val	G1y	Ser	Ile	Met	Pro	Pro	Phe	Thr	
		210	,				215					220	•				
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	Cys	His	Ser	Cys	Gly	Arg	Pro	Met	Gln	Gly	Pro	Ala	Ala	Pro	Asp	Val	
15	225			•		230					235					240	
20	ata	cac	gca	ccc	ggt	ccc	ctc	gac	gtt	gtc	acc	cct	gça	ctt	ggc	atg	768
	Ile	His	Ala	Pro	G1y	Pro	Leu	Asp	Val	Val	Thr	Pro	Ala	Leu	G1 y	Met	
		•			245			•		250					255		
25					<del></del>								•		200		
	ggc	ctc	ggt	ctc	tct	gac	cat	ggc	gct	gcc	gag	ctc	aga	cag	aaa	ctt	816
30					Ser												
	•			260				,	265					270		200	
									200					210			٠
35	886			go+			<b>770</b>	<b>600</b>	222	~~+	a	222	-++				064
																ccc	864
40	Oly	rne		vsħ	His	GIU	'ASP		1111	GIY	ser		٠	AST	Leu	Pro	
40			275					280					285				
<b>4</b> 5					agt										_		912
	Pro	G1y	Pro	Leu	Ser	Ala	Ala	Ala	Phe	Glu	Ser	Ala	Pro	Gly	Met	Ser	
		290					295					300					
50					•												
	gcc	gtc	gaa	gaa	ctc	aag	ctg	ctc	aag	gcc	cag	gtc	cag	gat	gtc	gct	960
55	Ala	Val	Glu	G1u	Leu	Lys	Leu	Leu	Lys	Ala	G1n	Val	G1n	^qzA	Val	Ala	

5	305			-		310					315				•	320	
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10	Arg	Val	Cys	Lys	Ala	Val	Ala	Glu	Gly	Asp	Leu	Ser	Gln	Lys	Ile	Thr.	
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15																	
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	Val	Pro	Val	Gln	Gly	Pro	Val	Met	Val	Gln	Leu	Lys	Asp	Val	Ile	Asn	
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										•		• •					
25	acc	atg	gtc	gat	aaa	cta	ggc	agg	ttt	gcg	cag	gag	gtc	act	cgt	gtc	1104
	Thr	Met	Val	Asp	Lys	Leu	G1y	Arg	Phe	Ala	Gln	G1u	Val	Thr	Arg	Val	
			355					360					365				
30			•							·							
	tcg	ctc	gaa	gtc	gga	act	gaa	ggc	cgg	ctc	ggt	ggt.	cag	gcc	att	gtt	1152
. 35	Ser		Glu	Val	Gly	Thr	Glu	Gly	Arg	Leu	G1y	Gly	Gln	Ala	Ile	Val	
-		370					375					380					
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		Asp	Va1	Arg	Gly	•	Trp	Ser	Glu	Leu		Thr	Val	Val.	Asn	_	
45 .	385					390					395	•				400	
											•		•				
		•						cag									1248
50	Leu	Ala	Ala			Thr	Ser	Gln	Val			Ile	Ala			Thr	
			•		405					410	•				415		•

	aag	gca	gto	gcc	aag	ggc	gat	ctc	tcc	aaa	caa	ato	ggc	gto	gat	gca	1296
5	Lys	Ala	Val	Ala	Lys	Gly	Asp	Leu	Ser	Lys	G1n	Ile	Gly	Val	Asp	Ala	
				420	)				425					430			
10																	
	aas	ggt	gaa	ata	ttg	gaa	ttg	aag	aat	acg	gtt	aat	acc	atg	gto	gtc	1344
	Lys	Gly	Glu	Ile	Leu	Glu	Leu	Lys	Asn	Thr	Val	Asn	Thr	Met	Val	Val	
15			435					440					445		•	•	
20	cgg	ttg	cgt	atg	ttt	gca	ggc	gaa	gtc	acc	cga	gtc	gcg	ctc	gat	gtc	1392
20					Phe												
		450					455					460					
25		•						•							•		
	ggc	agt	cgt	ggt	att	cta	99C	ggt	cag	got	tat	etc	cca	gat	atc	T2 T	1440
					Ile												1440
30	465			<b>02</b> ,		470	01,	<b>01</b> )	V-111	MAG	475		110	nsp	141		
	100					7,0					410					480	
35	aat	<b>**</b>	+														1.400
-					gag												1488
	GTA	val	irp	GIN	Glu		Ihr	Asp	Asn		Asn	Arg	Met	Cys		Asn	
40		•		٠	485		•	•		<b>4</b> 90					495		
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<b>1</b> 5					gtc												1536
	Leu	Thr	Asn	Gln	Val	Arg	Ser	Ile	Ala	Leu	Val	Thr	Thr	Ala	Val	Ala	
				500				•	<b>50</b> 5					510			
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5	Glu	Gly	Asp	Leu	Thr	Arg	Lys	Ile	G1u	Ile	Glu	Val	Glu	Gly	G1u	Met	
-																	

5			515	;				520					525	i			
10					aat Asn											acg	1632
	_	530				•	535				,	540			002	•••	
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	Phe	Ala	Ser	Glu	Val	Thr	Arg	Val	Ala	Leu	Glu	Val	Gly	Ser	Met	Gly	
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. 35	Asp	Leu	Thr	Arg	Asn	Val	Asn	Asn	Met	Ala	Ser	Asn	Leu	Thr	Asn	Gln	
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	Val	Arg	Ser	Ile	Ala	Lys	Val	Thr	Thr	Ala	Val	Ala	His	Gly	Asp	Leu	
45	٠		<b>5</b> 95				. •	600					605				
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50	Arg	Gln	Phe	Val	Glu	Val.	Asp	Val	G1n	Gly	Glu	Met	Leu	Met	Leu	Lys	
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5	Asn	Thr	· Val	Asn	Ser	Met	Val	Ala	Gln	Leu	Asp	Thr	Leu	Ala	Ser	- Glu	
	625	;				630	ı				635	;				640	
10									•								
	gtg	tcg	cgt	gtc	gcg	ctc	gag	gtc	ggt	atc	gag	ggt	cga	cto	ggt	gga	1968
	Val	Ser	Arg	Val	Ala	Leu	Glu	Val	Gly	Ile	Glu	Gly	Arg	Leu	Gly	Gly	
15					645					650			•		655		
	•																
20	cag	gct	gtg	gtt	cag	ggt	gtg	gag	ggt	gtg	tgg	aag	gtt	tta	acg	gac	2016
	Gln	Ala	Val	Val	G1n	G1y	Val	Glu	Gly	Val	Trp	Lys	۷al	Leu	Thr	Asp	
				660					665					670			
25	•					-											
	aat	gtc	aac	ttg	atg	gct	ctg	aat	ctg	acg	acc	caa	gtg	cgg	tct	att	2064
30	Asn	Val	Aşn	Leu	Met	Ala	Leu	Asn	Leu	Thr	Thr	Gln	Val	Arg	Ser	Ile	
•			675					680		_			685				
35											•						
-	gcg	gct	gtg	acg	act	gcc	gtg	gcg	cgt	ggt	gac	ctt	agc	aag	aat	atc	2112
	Ala	Ála	Val	Thr	Thr	Ala	Val	Ala	Arg	Gly	Asp	Leu	Ser	Lys	Asn	Ile	
40		690					<b>695</b>		•			700					
45	gat	gtc	gat	gtc	aag	ggc	gag	att	ttg	gat	ttg	aag	att	acg	gtc	aat	2160
	Asp	Val	Asp	Val	Lys	Gly	G1u	Ile	Leu	Asp	Leu	Lys	Ile	Thr	Val	Asn	•
	705					710	•				715					720	
50																	
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	Arg	Met	Thr	Asp	Ser	Leu	Arg	Ile	Phe	Ala	Ala	Glu	Val	Thr	Arg	Val	

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		gcg	cgc	gag	gtc	ggt	acg	ctc	gga	cga	ctc	ggc	gga	cag	gcg	ttt	gtt	2256
10		Ala	Arg	Glu	Val	Gly	Thr	Leu	Gly			Gly	Gly	Gln		Phe	Val	
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. •					885					890					895		
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					Lys												
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															•		
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			•		Ser			-					•				
55				•		•										•	

		930	)				935	5				940	)				
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	acı	g CCE	g ate	g aac	gge	att	att	ggc	ate	ace	gat	cto	acg	ctt	gat	acc	2880
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•	Gln	Asn	Leu	Asn	Leu	Phe	Tyr	Glu	Val	Asp	Pro	G1u	Ile	Pro	Asp	Gln.	
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	Va)	Ile	e G1)	/ Asp	Ser	Leu	Arg	Leu	Arg	Gln	Val	Ile	Thr	Asn	ı Lei	ı Val	
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				•													
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	Val Ser Leu Met	Asn Gly Gln Me	t Trp Val Glu Se	r Glu Val Gl	y Val
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	Gly Ser Arg Phe	Tyr Phe Thr Il	e Thr Ala Glu Il	e Ser Arg Pro	o Asn
	1155	116	0	1165	
15					
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20	Met Ala Gln Ser	Leu Gln Lys Va	l Ala Ile Tyr Ly:	s Glu Arg Thi	r Ile
	1170	1175	1180	)	
	•				
25	ttg ttt gtc gat	act ctg ggc gad	cgg tcg ggt gtg	gcg gag cgt	atc 3600
	Leu Phe Val Asp	Thr Leu Gly Asp	Arg Ser Gly Val	. Ala Glu Arg	Ile
30	1185	1190	1195	•	1200
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35	Glu Glu Leu Gln	Leu Arg Pro Phe	Yal Val Arg Asp	Ile Ser Gln	Val
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40			·		
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	Ala Asp Lys Ala	Lys Ile Pro Phe	Ile Asp Thr Val	Ile Val Asp	Ser
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	•	-			
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	•		Glu Leu Asp His		
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					Glu												
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									,					•			
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30					Thr								•				3300
· •				1300		110	Jei		305	VIG	1 7 1	nsp		1310	Leu	nia	
- 35				1300				4	1303			-		1310			
-	700	***	22+		-+-						-4-		4				2004
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40	GIU			vaı	Val	ASN	•		vai	Ala	Val			Leu	Glu	Lys	
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	ttt	ggt	cac	acg	gtt	cag	att	gcc	gag	aat	gga	cag	ttt	gcg	gtc	gac	4032
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	ant and atm	·				
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-	1410				Arg Thr Asp Leu	
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40	gtg age geg	2TC 222 CCC	ata eta con	ccc cac ggt		4017
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50	1 5 10 15														
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	\[ Va]	His	Gln	Leu	Ala	Lys	Ser	Leu	Leu	Leu	Ile	Ile	Asp	Asp	Ile	Leu	
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50									100					110			
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. ,			50					55					60				
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	·	<b>6</b> 5			•		70					75					80
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50				115					120								

#### Claims

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- A transformed cell in which a polynucleotide having a nucleotide sequence encoding an amino acid sequence of an osmosensing histidine kinase having no transmembrane region has been introduced in a functional form into a cell deficient in at least one hybrid-sensor kinase.
- 2. A transformed cell according to claim 1, wherein the polynucleotide having a nucleotide sequence encoding an amino acid sequence of an osmosensing histidine kinase having no transmembrane region is a polynucleotide complementing the deficiency in hybrid-sensor kinase in the cell deficient in at least one hybrid-sensor kinase in which the polynucleotide has been introduced.
- 3. A transformed cell according to claim 1 or 2, wherein the cell is a microorganism.
- 4. A transformed cell according to claim 3, wherein the microorganism is budding yeast.
- 5. A transformed cell according to any one of claims 1 to 4, wherein the osmosensing histidine kinase having no transmembrane region is an osmosensing histidine kinase having no transmembrane region and having a mutation which confers resistance to any of a dicarboxyimide antifungal compound, an aromatic hydrocarbon antifungal compound and a phenylpyrrole antifungal compound to the cell.
- A transformed cell according to claim 5, wherein the osmosensing histidine kinase having no transmembrane region is a histidine kinase having the amino acid sequence represented by SEQ ID NO: 13.
- A transformed cell according to any one of claims 1 to 5, wherein the osmosensing histidine kinase having no transmembrane region is an osmosensing histidine kinase which is derived from a plant-pathogenic filamentous fungus and has no transmembrane region.
  - 8. A transformed cell according to any one of claims 1 to 5 and 7, wherein the osmosensing histidine kinase having no transmembrane region is an osmosensing histidine kinase which is derived from Botryotinia fuckeliana, Magnaporthe grisea, Fusarium oxysporum, Mycospharella tritici, Thanatephorus cucumeris or Phytophthora infestans, and has no transmembrane region.
  - A transformed cell according to claim 1, wherein the osmosensing histidine kinase having no transmembrane region is an osmosensing histidine kinase having no transmembrane region which has an amino acid sequence represented by SEQ ID NO: 1, SEQ ID NO: 16, SEQ ID NO: 41, SEQ ID NO: 55, SEQ ID NO: 68 or SEQ ID NO: 90.
  - 10. A transformed cell according to claim 1, wherein the nucleotide sequence encoding an amino acid sequence of the osmosensing histidine kinase having no transmembrane region is a nucleotide sequence represented by SEQ ID NO: 2, SEQ ID NO: 17, SEQ ID NO: 42, SEQ ID NO: 56 or SEQ ID NO: 69.
  - 11. A method of assaying the antifungal activity of a substance, which comprises:
    - a first step of culturing a transformed cell as defined in any one of claims 1 to 10 in the presence of a test substance;
    - a second step of measuring an amount of intracellular signal transduction from the osmosensing histidine kinase having no transmembrane region expressed in the transformed cell cultured in the first step or an index value having the correlation therewith; and
    - a third step of assessing the antifungal activity of the test substance based on a difference between an amount of intracellular signal transduction or an index value having the correlation therewith measured in the second step and a control.
  - 12. A method according to claim 11, wherein the amount of intracellular signal transduction from the osmosensing histidine kinase having no transmembrane region or the index value having the correlation therewith is an amount of growth of the transformed cell.
  - 13. A method of identifying an antifungal compound, which comprises selecting an antifungal compound based on the antifungal activity assessed in the assaying method as defined in claim 11.

14. An antifungal compound selected by a method as defined in claim 13.

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- A method of killing a fungus, which comprises identifying an antifungal compound by a method as defined in claim
   and contacting the fungus with the identified antifungal compound.
- 16. An osmosensing histidine kinase having no transmembrane region, wherein the kinase is derived from a plant-pathogenic filamentous fungus.
- 17. An osmosensing histidine kinase having no transmembrane region, which has an amino acid sequence selected from the group consisting of:
  - (a) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which has an amino acid sequence homology of 95% or more to the amino acid sequence represented by any of SEQ ID NOs: 41, 55 and 68;
  - (b) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Fusarium oxysporum-derived cDNA as a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 52 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 53 as primers;
  - (c) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Mycospharella tritici-derived cDNA as a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 64 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 65 as primers;
  - (d) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Thanapethorus cucumeris-derived cDNA as a template andusing an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 85 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 86 as primers;
  - (e) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is derived from Phytophthora infestans and has the amino acid sequence represented by SEQ ID NO: 90;
  - (f) the amino acid sequence represented by SEQ ID NO: 41;
  - (g) the amino acid sequence represented by SEQ ID NO: 55; and
  - (h) the amino acid sequence represented by SEQ ID NO: 68.
- 18. An osmosensing histidine kinase having no transmembrane region, which has an amino acid sequence represented by SEQ ID NO: 41, SEQ ID NO: 55 or SEQ ID NO: 68.
- 19. A polynucleotide having a nucleotide sequence encoding an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, wherein the kinase is derived from a plant-pathogenic filamentous fungus.
- A polynucleotide having a nucleotide sequence encoding an amino acid sequence selected from the group consisting of:
  - (a) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which has an amino acid sequence homology of 95% or more to the amino acid sequence represented by any of SEQ ID NOs: 41, 55 and 68;
  - (b) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Fusarium oxysporum-derived cDNA as a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 52 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 53 as primers;
  - (c) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Mycospharella tritici-derived cDNA as a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 64 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 65 as primers;
  - (d) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is encoded by a DNA amplified by a polymerase chain reaction using a Thanapethorus cucumeris-derivedcD-NAas a template and using an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 85 and an oligonucleotide having the nucleotide sequence represented by SEQ ID NO: 86 as primers;
  - (e) an amino acid sequence of an osmosensing histidine kinase having no transmembrane region, which is derived from Phytophthora infestans and has the amino acid sequence represented by SEQ ID NO: 90;

- (f) the amino acid sequence represented by SEQ ID NO: 41;
- (g) the amino acid sequence represented by SEQ ID NO: 55; and
- (h) the amino acid sequence represented by SEQ ID NO: 68.

- <sup>5</sup> 21. A polynucleotide having a nucleotide sequence represented by SEQ ID NO: 42, SEQ ID NO: 56 or SEQ ID NO: 69.
  - 22. A method of obtaining a polynucleotide having a nucleotide sequence encoding an amino acid sequence of osmosensing histidine kinase which is derived from a plant-pathogenic filamentous fungus and has no transmembrane region, which comprises a step of amplifying a desired polynucleotide by Polymerase Chain Reaction using an oligonucleotide having a nucleotide sequence represented by any of SEQ ID NOs: 30 to 40, 52, 53, 64, 65, 85 and 86 as primers, and a step of recovering the amplified desired polynucleotide.
  - 23. An oligonucleotide which comprises a nucleotide sequence represented by any of SEQ ID NOs: 30 to 40, 52, 53, 64, 65, 85 and 86.